Delta Doping of Ferromagnetism in Antiferromagnetic Manganite Superlattices

T. S. Santos, B. J. Kirby, S. Kumar, S. J. May, J. A. Borchers, B. B. Maranville, J. Zarestky, S. G. E. te Velthuis, J. van den Brink, and A. Bhattacharya

1Center for Nanoscale Materials, Argonne National Laboratory, Argonne, Illinois 60439, USA
2NIST Center for Neutron Research, NIST, Gaithersburg, Maryland 20899, USA
3Institute for Theoretical Solid State Physics, IFW Dresden, 01171 Dresden, Germany
4Department of Materials Science and Engineering, Drexel University, Philadelphia, Pennsylvania 19104, USA
5Materials Science Division, Argonne National Laboratory, Argonne, Illinois 60439, USA
6Ames Laboratory and Department of Physics and Astronomy, Iowa State University, Ames, Iowa 50011, USA

(Received 10 July 2011; published 12 October 2011)

We demonstrate that delta doping can be used to create a dimensionally confined region of metallic ferromagnetism in an antiferromagnetic (AFM) manganite host, without introducing any explicit disorder due to dopants or frustration of spins. Theoretical consideration of these additional carriers shows that they cause a local enhancement of ferromagnetic double exchange with respect to AFM superexchange, resulting in local canting of the AFM spins. This leads to a highly modulated magnetization, as measured by polarized neutron reflectometry. The spatial modulation of the canting is related to the spreading of charge from the doped layer and establishes a fundamental length scale for charge transfer, transformation of orbital occupancy, and magnetic order in these manganites. Furthermore, we confirm the existence of the canted, AFM state as was predicted by de Gennes [Phys. Rev. 118, 141 (1960)] but had remained elusive.

DOI: 10.1103/PhysRevLett.107.167202 PACS numbers: 75.70.Cn, 61.05.fj, 73.21.Cd, 75.47.Lx

In semiconductors, delta doping [1] has led to record high mobilities in two-dimensional electron gases, spawning a number of fundamental discoveries and many novel applications. In the complex oxides, similar doping strategies may in principle be used to create two-dimensional analogs of the many collective phases found in these materials [2,3]. In this work, using antiferromagnetic (AFM) La$_{1-x}$Sr$_x$MnO$_3$ as the host material, we have devised a delta-doping strategy for locally tailoring the strength of the ferromagnetic (FM) double-exchange interactions relative to the AFM superexchange, creating a quasi-two-dimensional region of enhanced moment. We do this without introducing a random disorder potential due to dopants or the frustration inherent to some AFM-FM interfaces. Our approach is unique to systems where double-exchange (DE) and superexchange (SE) interactions compete and cannot be realized in conventional semiconductors or metals.

In the seminal work of de Gennes [4], it was recognized that when the insulating AFM parent compound of a manganite is doped with carriers, competition between DE and SE causes the AFM spins to cant, in proportion to the doping. In real materials, additional effects such as the Jahn-Teller (JT) interactions, on-site Coulomb repulsions (Mott-Hubbard effects), and orbital ordering instabilities, all favor localizing the carriers and compete with DE. As a result, the transition between an AFM insulator such as LaMnO$_3$ to a FM metal such as L$m$O$_{100}$Sr$_{20}$MnO$_3$ does not follow de Gennes’ phase diagram but rather takes place through a mixed-phase region for intermediate values of $x$ [5]. A tendency toward phase separation is believed to be a fundamental property of the double-exchange manganites [6] and may thwart the appearance of a homogeneous, canted AFM phase [7].

The situation is quite different if one starts with La$_{1-x}$Sr$_x$MnO$_3$ for $x > 0.5$, which is also AFM. The Mn $e_g$ electrons can go into one of two partially filled bands made of degenerate $d_{3z^2-r^2}$ and $d_{r^2}$ orbitals. For 0.5 < $x$ < 0.7, this degeneracy is removed by the formation of a highly anisotropic $A$-type AFM order (planes of FM spins that are mutually AFM), with the electrons mainly occupying the $d_{3z^2-r^2}$ orbital states [8]. The in-plane transport is nearly metallic due to DE. However, the out-of-plane resistivity is orders of magnitude higher since the $d_{3z^2-r^2}$ states are unoccupied and SE dominates in this direction [9]. Upon doping with electrons toward $x < 0.5$, the $d_{3z^2-r^2}$ orbitals begin to be occupied, DE begins to act in the out-of-plane direction, and the material transforms into a three-dimensional FM metal. We show in this work that by delta doping an AFM manganite superlattice (SL) with electrons, we locally enhance the DE, which cant the spins away from AFM alignment, resulting in a highly modulated magnetization. Our theoretical investigation shows that a JT distortion is instrumental in stabilizing this canted spin structure lying in the midst of the crossover from a 2D antiferromagnet to a 3D ferromagnet near $x = 0.5$. We synthesized cation-ordered analogs of La$_{0.5}$Sr$_{0.5}$MnO$_3$ by alternating single unit cell layers of LaMnO$_3$ (LMO) and SrMnO$_3$ (SMO) with atomic layer precision onto SrTiO$_3$ substrates using ozone-assisted molecular beam epitaxy. Neutron diffraction measurements on these (LaMnO$_3$)$_1$/SrMnO$_3$$_1$ superlattices confirmed...
the $A$-type AFM phase, and magnetometry measurements using a superconducting quantum interference device (SQUID) showed a net FM moment near zero at low field [10]. Here, we present delta-doped SLs with nominal compositions $x = 0.44$ and $0.47$. The $x = 0.44$ SL was made by alternating unit cell layers of LaMnO$_3$ and SrMnO$_3$ and inserting an additional LaMnO$_3$ layer for every four LaMnO$_3$/SrMnO$_3$ bilayers [11]. This sequence was repeated 9 times (9 supercells), to form the superlattice [(SMO1/LMO1) $\times 4$, LMO1] $\times 9$. A La$_{0.56}$Sr$_{0.44}$MnO$_3$ alloy film with a thickness of 81 uc was made for direct comparison. The $x = 0.47$ SL was made by inserting an extra LaMnO$_3$ layer after every 9 LaMnO$_3$/SrMnO$_3$ bilayers, repeated 4 times: [(SMO1/LMO1) $\times 4$, LMO1, (SMO1/LMO1) $\times 5$] $\times 4$. Structural characterization of the SLs is discussed elsewhere [10,11]. By periodically inserting the extra LaMnO$_3$ unit cell layer, we have delta doped the MnO$_2$ planes in the vicinity of the inserted LaMnO$_3$ unit cell with electrons. Our investigation of these SLs by neutron scattering techniques was aimed at determining the influence of these delta-doped charges on the $A$-type spin structure.

We performed neutron diffraction measurements to probe the spin structure in the $x = 0.44$ SL and alloy. The $A$-type order is manifested by a structurally forbidden (0 0 1/2) Bragg peak, shown in Fig. 1(a), signifying an AFM spin alignment with a periodicity of $2c$ normal to the film plane ($c$ is the out-of-plane lattice parameter). From the full width at half maximum of the peak, a magnetic coherence length of $\sim 28.5$ nm was determined, nearly the entire film thickness (30.9 nm). The coexistence of both a (0 0 1/2) neutron diffraction peak and a net FM moment from $M(H)$ data [11] suggests a canted AFM structure, for which the spins within a single plane are ferromagnetically aligned and the spins on adjacent planes are canted at a relative angle $\theta_c < 180^\circ$, where $\theta_c/2$ is the angle between each spin and the applied magnetic field direction. The temperature ($T$) dependence of the peak intensity in Fig. 1(b) indicates that the $A$-type order in the SL disappears at $\sim 260$ K.

To determine the depth-dependent magnetization profile, we performed polarized neutron reflectometry (PNR) measurements, which detect variations of in-plane magnetization as a function of sample depth along the surface normal. The non-spin-flip reflectivities $R_{++}^\perp$ and $R_{--}^\perp$, shown in Fig. 2(a) for the $x = 0.44$ SL, are sensitive to the component of magnetization parallel to the applied field direction. Because the nuclear scattering length densities of SMO and LMO are nearly identical, only magnetic variations in the layer profile are detectable. The key feature of interest here is the Bragg peak at $q = 0.18$ A$^{-1}$, indicating that the magnetization is in fact modulated with a periodicity that matches the supercell periodicity of the SL.

To fit the PNR data by using the CO_REFINE routine [12], a model consisting of 9 supercells was used, where each supercell consisted of a high moment sublayer and a low moment sublayer. The sublayer thicknesses, magnetic roughnesses, and magnetic scattering length densities (a quantity directly proportional to the magnetization) of the seven interior supercells were fit while constrained to be identical, whereas the topmost and bottommost supercells could vary independently. For these data and model, there are several sets of possible fit parameters that yield the same lowest $\chi^2$ value to within 1%. Two such profiles are shown in Fig. 2(b). Even though the highest and lowest moments in the modulation period, $M_{\text{max}}$ and $M_{\text{min}}$, respectively, can each vary by as much as 0.25 $\mu_B$, we can conclude with certainty across all sets of fit parameters that (i) the period of the magnetic modulation matches the structural supercell to within 0.1 nm, (ii) the high moment sublayer contains the extra LMO layer, and (iii) the magnitude of the modulation is quite large, with $M_{\text{max}}/M_{\text{min}} = 3 \pm 0.3$.

The $x = 0.47$ SL has a thicker supercell with one extra LMO layer for every 19 uc, such that two Bragg peaks are detectable, as shown in the PNR spectra in Fig. 2(c). With this additional information on the magnetic modulation compared to the $x = 0.44$ SL data set, the fitting routine produces a single set of best fit parameters at the lowest $\chi^2$ for the $x = 0.47$ SL. A model similar to the one previously described, containing alternating high moment and low moment sublayers, was used to fit these data. The moment is again highly modulated, alternating between a high moment region across 6 uc [the full width at half maximum of the peaks in Fig. 2(d)] containing the extra LMO unit cell and having $M_{\text{max}} = 2.2 \mu_B$ and a low moment region spanning 13 uc having $M_{\text{min}} = 0.7 \mu_B$. The gradual transition between $M_{\text{max}}$ and $M_{\text{min}}$ occurs over 4 uc, and, as we shall show, this occurs by varying the canting angle. By using $M_{\text{tot}} = (4 - x) \mu_B$ as the magnitude of the moment of a Mn$^{3+}$/4+ spin and $M = M_{\text{tot}} \cos(\theta_c/2)$, the measured modulation of $M$ corresponds to a modulation of canting angle between $\theta_c = 103^\circ$ and $157^\circ$. These fit parameters also work for the $x = 0.44$ SL, where the high moment region extends across 6 uc with $M_{\text{max}} = 2.2 \mu_B$ and the low moment region extends across 3 uc with $M_{\text{min}} = 0.7 \mu_B$, as shown in Fig. 2(d).
are the fit to the data. The inset shows the PNR of the signifying that the magnetization is modulated. The solid lines 675 mT same data multiplied by 66. The Bragg peaks are more apparent in the inset, which shows the profiles as a function of film depth from the fit to the PNR spectra (green) layers in the SL and the SrTiO$_3$ substrate (gray) are shown. (c) PNR for the x = 0.47 SL with H = 815 mT and T = 120 K. The Bragg peaks are more apparent in the inset, which shows the same data multiplied by $q^2$ to compensate for Fresnel decay. (d) The magnetic depth profile of the x = 0.47 SL from the best fit. shown in Fig. 2(b) (black line). In comparison, the PNR measurement of the x = 0.44 alloy film [inset in Fig. 2(a)] shows a net moment without any modulation (no Bragg peak). Thus, the spin canting in the alloy is uniform, and

![FIG. 2](color online). (a) The non-spin-flip channels ($R^{++}$ and $R^{--}$) of the PNR for the x = 0.44 SL at T = 120 K and with H = 675 mT applied in the plane of the film have a Bragg peak signifying that the magnetization is modulated. The solid lines are the fit to the data. The inset shows the PNR of the x = 0.44 alloy film having no Bragg peak. (b) Two possible magnetic profiles as a function of film depth from the fit to the PNR spectra of the x = 0.44 SL. The locations of the LMO (pink) and SMO (green) layers in the SL and the SrTiO$_3$ substrate (gray) are shown. (c) PNR for the x = 0.47 SL with H = 815 mT and T = 120 K. The Bragg peaks are more apparent in the inset, which shows the same data multiplied by $q^2$ to compensate for Fresnel decay. (d) The magnetic depth profile of the x = 0.47 SL from the best fit. the modulated magnetization in the x = 0.44 and 0.47 SLs occurs due to the delta doping.

To investigate this canted spin structure further, we carried out diffraction measurements with polarized neutrons and polarization analysis in the vicinity of the AFM peak for the x = 0.44 SL, as shown in Fig. 3(a). The peak in spin-flip intensities $I^{+-}$ and $I^{-+}$ at q = 2$\pi$/(2c) = 0.82 Å$^{-1}$ results from the canted AFM spins with each sublattice having a component of magnetization pointing perpendicular to the applied field direction, with a period of 2c. Moreover, the superlattice peak at q = 0.64 Å$^{-1}$ signifies that the canted spin structure is modulated with a period of 9 uc, consistent with the supercell period and the periodicity of the FM component revealed by the low q PNR measurements [13]. These peaks in $I^{+-}$ and $I^{-+}$ directly confirm the canted, modulated spin structure. Figure 3(b) shows the canted, modulated spin structure as deduced by the PNR and diffraction measurements.

In order to explore theoretically this canted spin state, we consider the standard two-orbital model for manganites [7] with the Hamiltonian $H = H_{DE} + H_{SE} + H_{JT}$ [11]. The double-exchange term $H_{DE}$, which arises from a combination of kinetic energy and strong Hund’s rule coupling, favors the FM state. The superexchange term $H_{SE}$ is the Heisenberg interaction between $t_{2g}$ spins favoring AFM coupling. The Jahn-Teller term $H_{JT}$ represents the coupling of the $e_g$ electrons to the lattice distortions.

Before understanding the canted magnetic states, it is essential to understand the stability of A-type AFM order near x = 0.5 in manganites. It is an outcome of a competition between the ferromagnetic
From a simple energy minimization standpoint, this causes occupancy of the $d_{3z^2-r^2}$ orbitals to decrease, which in turn causes $J_F$ to decrease further, making the spins cant even more. The cost in elastic energy in creating the lattice distortion with changes in orbital occupancy (the $c$ axis shrinks with lower $p_z$) prevents the transition from being a step function.

In order to test this explicitly, we minimize the total energy of the Hamiltonian for variational parameters $\theta_z$ and JT distortions, hence allowing for this feedback effect. At a fixed $x$, JT distortions lead to a reduction in electronic energy, but there is also an elastic cost associated with making these distortions [11]. The optimum value of the distortion is selected from the above competition. This generates an $x$ dependence on the magnitude of the distortions and therefore an additional $x$ dependence on the values of $J_F$. The resulting optimum angle $\theta_z$ is plotted in Fig. 4(b) for a few values of $J_S$. The sharp change in $\theta_z$ from $180^\circ$ to $0^\circ$ for $J_S = 0.10, 0.11$ in the calculation does not allow for layers at intermediate $x$ with $0^\circ < \theta_z < 180^\circ$. This implies that it is not possible to stabilize a finite canting angle for these parameters. However, for $J_S \geq 0.12$ and for $x < 0.5$, the variation in $\theta_z$ with $x$ is more gradual and layers with a finite canting angle are stable. Therefore, a small variation in the doping level can lead to significant changes in the canting angle and thus the net FM moment. The large modulation in moment as a function of depth in these delta-doped SLs is a realization of this scenario. Attempts to realize the canted AFM state near $x = 0$ as proposed by de Gennes and near $x = 0.5$ have typically been obscured by phase segregation into FM and AFM domains [14], which is mitigated in our digital-synthesis approach of charge doping without disorder. We note that a modulation in orbital occupancy is expected to accompany this modulated spin state [15].

In conclusion, we have employed delta doping to tailor the magnetic structure following the ideas of de Gennes. Localized regions of enhanced FM exchange were created in an otherwise AFM structure, where the transition between the high moment region and the low moment region occurs via a continuous variation of the canting angle that is governed by the spreading of charge near the delta-doped layer. Furthermore, the PNR technique enables us to determine a length scale of $6 \text{ uc}$ for the region of enhanced moment that is induced by the delta-doped layer. The result that the SL magnetization is highly modulated is evidence that the doped charges are not completely delocalized over the entire SL. Charge spreading out to $3 \text{ uc}$ from both sides of the delta-doped layer is consistent with previous theoretical and experimental works that had inferred a length scale for charge spreading via more indirect means [16,17]. We believe this to signify a fundamental length scale for the spreading of charge normal to the layers.

We are grateful to Chuck Majkrzak for helpful suggestions. Work at Argonne and use of the Center for Nanoscale Materials was supported by the U.S. Department of Energy, Office of Science, Office of Basic Energy Sciences, under Contract No. DE-AC02-06CH11357. A portion of this research at Oak Ridge National Laboratory’s High Flux Isotope Reactor was sponsored by the Scientific User Facilities Division, Office of Basic Energy Sciences, U.S. Department of Energy. We
acknowledge the support of the National Institute of Standards and Technology, U.S. Department of Commerce, in providing the neutron research facilities used in this work. T. S. acknowledges support from the L’Oreal USA Fellowship for Women in Science.

*anand@anl.gov

[13] The SL periodicity (9c) is found from the distance between the two peaks in Fig. 3(a): 0.82 Å⁻¹ − 0.64 Å⁻¹ = 0.18 Å⁻¹ = 2π/(9c).