Electronic Confinement and Ordering Instabilities in Colossal Magnetoresistive Bilayer Manganites


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We present angle-resolved photoemission studies of (La$_{1-x}$Pr$_x$)$_2$Sr$_2$Mn$_2$O$_7$ with $x = 0.4$ and $z = 0.1$, $0.2$, and $0.4$ along with density functional theory calculations and x-ray scattering data. Our results show that the bilayer splitting in the ferromagnetic metallic phase of these materials is small, if not completely absent. The charge carriers are therefore confined to a single MnO$_2$ layer, which in turn results in a strongly nested Fermi surface. In addition to this, the spectral function also displays clear signatures of an electronic ordering instability well below the Fermi level. The increase of the corresponding interaction strength with $z$ and its magnitude of ~400 meV make the coupling to a bare phonon highly unlikely. Instead we conclude that fluctuating order, involving electronic and lattice degrees of freedom, causes the observed renormalization of the spectral features.

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An important issue in the field of correlated electron physics concerns the relationship of collective charge carrier dynamics and electronic ordering tendencies [1]. In the limiting case where the interactions between the electrons are weak and the kinetic energy dominates, the resulting many-particle system can be described as a metallic Fermi liquid of noninteracting quasiparticles. This changes dramatically if the interactions between the electrons are strong and the dominant role of the kinetic energy is lost. In this limit, the charge carriers can crystallize, resulting in insulating behavior and static spatial modulations of the electronic system. In contrast to the Fermi liquid, these electronic crystals typically break the symmetry of the underlying nuclear lattice.

A most intriguing case might be realized just in between these two limits, where neither the interactions between the electrons nor the kinetic energy is dominating. In these cases, electronic liquid crystals may emerge, where the electrons crystallize to some extent, while still forming a conductive phase [2]. Electronic liquid crystals, and more specifically nematic electronic order, are currently discussed intensively in relation to the unconventional superconductivity and the pseudogap phase of the high-temperature superconducting cuprates [3–5]. There are also clear indications for such an ordering in some of the new iron pnictide superconductors [6].

The observed proximity of metallic and ordered phases in doped manganites shows that also in these materials ordering tendencies and kinetic energy both are important and competing [7,8]. It is also widely believed that this competition lies at the heart of the famous colossal magnetoresistance (CMR) effect, but the detailed physical mechanisms driving the CMR still remain to be fully understood. In the bilayer CMR manganites La$_{1.2}$Sr$_{1.3}$Mn$_2$O$_7$ ($x = 0.4$), the competition between electronic ordering tendencies and charge delocalization is particularly prominent: First, the growth of ordered regions with cooling in the paramagnetic insulating phase is observed in scattering experiments [9]. Then, when the transition into the ferromagnetic metallic (FMM) phase is reached at $T_C$, these ordered regions melt and the CMR occurs [9,10]. Correspondingly, unconventional electronic properties of the FMM phase were indeed observed in previous angle-resolved photoemission spectroscopy (ARPES) studies, including a strongly doping and temperature dependent pseudogap [11–13].

In this Letter, we present a study of (La$_{1-x}$Pr$_x$)$_2$Sr$_{1.3}$Mn$_2$O$_7$ (LPSMO) with a fixed hole concentration of $x = 0.4$. Previous macroscopic studies of these samples showed that the FMM phase is strongly destabilized with increasing Pr concentration $z$ [10]. The reduction of $T_C$ with increasing $z$ indicates that the chemical pressure due to the smaller Pr affects the balance between kinetic energy and electronic ordering tendencies. Here, we apply ARPES to investigate how altering this balance changes the electronic structure and the charge dynamics of LPSMO. ARPES provides direct access to the single-particle spectral function $\mathcal{A}(\mathbf{k}, \omega)$ [14] and, in combination with density functional theory calculations and x-ray diffraction data, enables us to reveal the effects of fluctuating electronic order on the charge dynamics in the FMM phase of LPSMO.

The studied LPSMO single crystals were grown by the traveling floating zone method. It was found that a small
amount of Pr enables growth of single crystals of very high quality as confirmed by x-ray diffraction, magnetization, and thermal transport measurements. The ARPES measurements were performed using the I\(^3\) station at the beam line UE112 of the synchrotron facility BESSYII, which provides a beamspot of about 100 \(\mu m \times 100 \mu m\). All presented ARPES spectra were measured with 55 eV excitation energy at around 23 K, if not stated otherwise. The diffuse x-ray diffraction experiment was performed at the high-energy beam line BW5 at HASYLAB in Hamburg. At the used photon energy of 100 keV, the penetration depth is of the order of 1 mm, guaranteeing the detection of true bulk properties. The density functional theory calculations in the local spin density approximation (LSDA) and the subsequent tight-binding (TB) analysis were done in the code FPLO 8.5 [15]. All calculations for the hole doping of \(x = 0.4\) were done in the virtual crystal approximation.

Figures 1(a)–1(c) show the measured Fermi surface (FS) of LPSMO with \(z = 0.1\). Very similar results were obtained for \(x = 0.4\). These data were taken deep inside the FMM phase (\(T_C = 100\) K) and the polarization was chosen to enhance the spectral weight at the Fermi level, as demonstrated in Fig. 1(e). The energy distribution curves (EDCs) in Fig. 1(e) also show that the spectral weight at the Fermi energy \(E_F\) is small, which is in line with the small Drude weight observed for these compounds [16]. The measured remnant FS formed by these states consists of parallel straight sections over large regions of \(k\) space, which agrees with previous ARPES studies. This well-nested FS already indicates the presence of electronic ordering instabilities.

According to LSDA, there is a sizable hybridization between the two planes within a bilayer, which is reflected by the large interplane hopping \(t_{\text{perp}} = 0.7\) eV obtained from a TB Wannier analysis. This large \(t_{\text{perp}}\) leads to a considerable splitting of bonding and antibonding bands, the so-called bilayer splitting (BLS). Along the \(M-X-M\) direction, for instance, the calculated BLS at \(E_F\) is about \(0.1 \times 2\pi/a\). As can be seen in Fig. 1(a), this does not agree with experiment, where such a big BLS is not observed. In fact, the data displayed in Fig. 1(f) clearly exclude the presence of a large BLS in LPSMO: even though the momentum distribution curves (MDCs) close to \(E_F\) are broad, their symmetric shape clearly shows that the BLS in the real system is much smaller, if not completely absent. This result was reproduced with different polarizations and in different regions in \(k\) space, excluding possible matrix element effects. The reduction of the BLS observed here is in perfect agreement with the results of Jozwiak et al., who also discovered a dramatic reduction of the BLS for the Pr-free compound between \(x = 0.36\) and 0.4 [17]. In addition, very similar FS were observed in single-layer manganites, where a BLS does not exist [18].

Within our TB model it is possible to set \(t_{\text{perp}} = 0\) eV, i.e., to switch off the bilayer hybridization and to eliminate the BLS. As can be seen in Fig. 1(b), this gives a much better agreement with experiment, further supporting the absence of the BLS. The FS of the real material is therefore very close to the independent particle reference given by this TB model. Importantly, the TB results given in Fig. 1(b) imply that setting \(t_{\text{perp}} = 0\) eV yields a strongly nested FS with large parallel sections, which is...
therefore also investigated

In order to study possible static ordering in our samples, we performed elastic high-energy x-ray diffraction (HEXRD) studies. Figure 3 shows the data obtained for samples with \( z = 0.2 \) and \( z = 0.4 \) at various temperatures around \( T_C \). Referring to the space group 14/mmm, weak intensity was detected at the symmetry forbidden (401) reflection due to a small amount of stacking faults within the probed sample volume.

More importantly, strong diffuse scattering intensity was observed around the (401). At room temperature, only broad diffuse scattering is observed (not shown) which has been attributed to uncorrelated polaronic lattice distortions [9]. Upon cooling down towards \( T_C \), superlattice peaks centered at \( (4 \pm 0.25, 0, 1) \) emerge out of this broad
by HEXRD agrees perfectly with the nesting vector $q$ for samples. Similar diffuse scattering results were obtained for Pr doping below $T_C$. The ordered regions melt upon entering the FMM phase.

Diffuse scattering, signaling the growth of ordered regions within the disordered majority phase. With entering the FMM phase at $T_C$, these superlattice peaks vanish abruptly and only diffuse scattering remains in the FMM phase. We also verified the absence of static CE order below $T_C$ in our samples. Similar diffuse scattering results were obtained for $La_{1.2}Sr_{1.8}Mn_2O_7$, where the temperature dependence of the broad superlattice reflections was discussed in terms of ordered areas with a correlation length of about 27 Å, which melt upon entering the FMM phase [9]. Interestingly, the modulation vector of (0.25, 0, 0) observed by HEXRD agrees perfectly with the nesting vector $q_1$ defined by the FS topology. This agreement between the FS topology, the backbending, and the measured ordering strongly argues in favor of ARPES representing bulk properties.

We note that the modulation vector (0.25, 0, 0) found for LPSMO differs from the (0.3, 0, 0) observed for $La_{1.2}Sr_{1.8}Mn_2O_7$ [9]. The reason for this difference is currently unclear. However, we can already exclude changes in the doping level caused by Pr, because the modulation vector determined by HEXRD for LPSMO is the same for $z = 0.2$ and 0.4. In addition, the FSs measured for $z = 0.1$ and $z = 0.4$ show no sign of a difference in hole doping. This interesting point is the subject of current investigations, but plays no role for the arguments presented in this Letter.

To summarize, the main result of the present ARPES study is the observation of clear fingerprints of electronic order deep inside the FMM phase of LPSMO. Apart from the strongly nested FS, which has also been reported previously for other bilayer compounds [23], we identified the presence of a backbending anomaly in the measured $A(k, \omega)$. The excellent agreement between experiment and the presented model calculation establishes the coupling of the charge carriers to a collective excitation, which causes an inherent breaking of translational symmetry. The large coupling strength $V = 432$ eV found for $z = 0.1$ makes a coupling to a bare phonon, as in the case of a conventional charge density wave transition, highly unlikely. This is further supported by the observed destabilization of the FMM phase with Pr doping [24], which is consistent with the measured increase in gap size [see Fig. 2(d)] and points towards an electronic origin of the underlying ordering phenomenon. At the same time, HEXRD shows no evidence for static order in the FMM phase. We therefore conclude that fluctuating order exists in the FMM phase, which has a strong impact on the charge carrier dynamics. Such an order is expected to involve all electronic and lattice degrees of freedom, which is a peculiarity of the mixed valence manganites.

A likely and consistent scenario is strong fluctuations between a two-dimensional metallic phase and the ordered CE phase, which both favor the occupation of in-plane orbitals. In fact, evidence for CE fluctuations in the FMM phase of $La_{1.2}Sr_{1.8}Mn_2O_7$ was recently deduced from inelastic neutron scattering data [25]. In addition, the TB analysis presented above clearly indicates that a two-dimensional metallic phase of LPSMO is indeed unstable towards electronic order. The notion of fluctuating CE order in the FMM phase close to $x = 0.5$ is therefore well supported by experimental and theoretical results. In particular, this scenario naturally explains the observed coexistence of a metallic FS and the backbending anomaly reported here.

Finally, we note that the FMM phase exhibits typical features of an electronic liquid crystal: (i) proximity to static order (diffuse scattering, FS nesting), (ii) evidence for molten electronic order (backbending, no static order), and (iii) reduced electronic dimensionality (FS, BLS). This strongly motivates further studies of CMR materials in this context.

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FIG. 3 (color online). X-ray diffuse scattering data of $(La_{1-x}Pr_x)_{2.5}Sr_{1.8}Mn_2O_7$ with $z = 0.2$ and $z = 0.4$ for different temperatures around $T_C$. The weak symmetry forbidden (4, 0, 1) Bragg peak is due to a small amount of stacking faults. Therefore, the (401) intensities of the two samples cannot directly be compared. Above the Curie temperature of around 90 K ($z = 0.2$) and 45 K ($z = 0.4$) broad superstructure reflections at $(4 \pm 0.25, 0, 1)$ are observed. The data sets at 88 K ($z = 0.2$) and 45 K ($z = 0.4$) were taken just above $T_C$. The ordered regions melt upon entering the FMM phase.