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Distinct Nature of Static and Dynamic Magnetic Stripes in Cuprate Superconductors

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We present detailed neutron scattering studies of the static and dynamic stripes in an optimally doped high-temperature superconductor, La2CuO4+y. We observe that the dynamic stripes do not disperse towards the static stripes in the limit of vanishing energy transfer. Therefore, the dynamic stripes observed in neutron scattering experiments are not the Goldstone modes associated with the broken symmetry of the simultaneously observed static stripes, and the signals originate from different domains in the sample. These observations support real-space electronic phase separation in the crystal, where the static stripes in one phase are pinned versions of the dynamic stripes in the other, having slightly different periods. Our results explain earlier observations of unusual dispersions in underdoped La2−xSrxCuO4 (x = 0.07) and La2−xBa4CuO4 (x = 0.095).

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An imperative open question in materials physics is the nature of high-temperature superconductivity. Unlike conventional superconductors, where the Cooper pairing mechanism is well established [1], the pairing mechanism in high-temperature superconductors (HTS) still sparks controversy [2]. A comprehensive description of the electronic behavior inside HTS is indispensable to push this field of research onward. Hence, the magnetic structures which appear close to as well as inside the superconducting phase are still being studied intensively [3,4]. In many HTS compounds, experiments indicate a modulated magnetic structure, consistent with superconducting “stripes” of charge separated by magnetic regions as sketched in Fig. 1(a) [5]. Magnetic excitations, referred to as “dynamic stripes,” are found with similar periodicity and are therefore thought to be related to the Goldstone modes of the static stripes [6].

Here we present evidence that this model is incomplete for a family of HTS. We find that the dynamic stripes do not disperse towards the static stripes in the limit of vanishing energy transfer and interpret this in terms of electronic phase separation, where static and dynamic stripes populate different spatial regions of the HTS.

Compounds based on the La2CuO4 family were the first HTS to be discovered [7]. They become superconducting upon doping with electrons or holes, with a maximum critical temperature \( T_c \approx 40 \) K, whether the dopant is Sr (La2−xSrxCuO4, LSCO), Ba (La2−xBaxCuO4, LBCO), or O (La2CuO4+y, LCO + O). The generic crystal structure of these compounds is illustrated in Fig. 1(b). They consist of planes of CuO2 separated by layers of (La,Sr) or (La,Ba). Each Cu atom is at the center of an octahedron of oxygen atoms. At elevated temperatures, these materials are in the high-temperature tetragonal phase. Upon lowering the temperature, the crystals enter the low-temperature orthorhombic phase (LTO), where the oxygen octahedra tilt around the tetragonal a axes, leading to a change in lattice parameters, \( a < b \), and to possible twinning [8,9]; see Supplemental Material for details [10].

Since the first discovery, a multitude of HTS have been found in the cuprate family. The amplitude and period of the stripe order modulations vary strongly with the choice and amount of dopant, with static stripes being particularly pronounced in LCO + O [11].

The spin stripes can be measured using magnetic neutron scattering, where they are observed as pairs of intensity peaks at incommensurate (IC) wave vector transfers, e.g., at \( \mathbf{Q} = (1 + \delta_h, \delta_k, 0) \) and \( \mathbf{Q} = (1 - \delta_h, -\delta_k, 0) \) for stripes along the (110) direction; see Fig. 1(c). Here, the components of the scattering vector are given in terms of \( (2\pi/a, 2\pi/b, 2\pi/c) \), where \( a, b, \) and \( c \) are the orthorhombic lattice constants. The real-space modulation period is \( L \sim (2\pi/\delta)a \), and we refer to \( \delta \) as the incommensurability of the stripes. Typically, \( \delta_h \approx \delta_k \), indicating that the modulation is approximately along the Cu–O–Cu bonds.
(the [110] and [1^-110] directions), although variations have been reported, indicating a kink in the stripes after a number of unit cells [11,12].

Typically, stripes are observed not only at the above mentioned positions but also at $Q = (1 - \delta_h, \delta_k, 0)$ and $Q = (1 + \delta_h, -\delta_k, 0)$, giving rise to a quartet of peaks around the (100) position, as illustrated in Fig. 1(d). This indicates that the compound exhibits stripes (approximately) along both the [110] and [1^-110] directions, most likely by the stripes in adjacent layers alternating between the [110] and [1^-110] directions [13].

Inelastic neutron scattering has shown the presence of dynamic stripes, which at low energies have a similar modulation period as the static stripes [14]. The modulation period of the stripes is found to be almost constant up to energy transfers of $\Delta E \sim 10 - 15$ meV [15,16]. In the cuprates, an hourglass-shaped dispersion develops at higher energies [17].

The incommensurability of the stripes varies with doping. In the LSCO-type cuprates, $\delta$ increases linearly with doping and saturates at a maximal value of $\delta = 1/8$ [14]. In some cuprates, similar stripes of charge with half the modulation period have been observed using x-ray diffraction, validating the picture of magnetic and charge stripes in Fig. 1(a) [18–22]. However, the energy resolution of x rays is not sharp enough to distinguish between static and low energy dynamics stripes.

We have used elastic and inelastic scattering of low-energy neutrons to accurately measure the reciprocal space position of the static and dynamic stripes in highly oxygenated LCO + O in the LTO phase. The experiments were performed at the cold-neutron triple axis spectrometers FLEXX at Helmholtz-Zentrum Berlin (HZB), Berlin [23], and Three Axis Low Energy Spectrometer (ThALES) at Institut Laue-Langevin (ILL), Grenoble [24]. The elastic energy resolution in the ThALES experiment was 0.24 meV [full width at half maximum (FWHM)], while the $Q$ resolution was 0.05 r.l.u. (FWHM). For further details on the experiments, see Supplemental Material [10].

Figure 1(d) shows how we probe two of the four IC peaks in our neutron scattering experiments. The actual data for a series of scans are shown in Figs. 1(e) and 1(f) as
2D color plots. Figure 2 shows examples of the scans through the center of the peaks at 0 and 1.5 meV energy transfer, probing the static and dynamic stripes, respectively. The inset illustrates the direction of the scans in reciprocal space.

To eliminate errors from minor misalignments, we determine the incommensurability along $k$, $\delta_k$, as half the distance between the peak centers. In Fig. 3, we display $\delta_k$ for all energy transfers probed in the experiment at two temperatures. As expected, the dynamic stripes appear at the same reciprocal space position in the normal phase (45 K) as in the SC phase (2 K) (within the instrument resolution), whereas the static stripes are present only at a low temperature. The elastic stripes are found to be rotated by $7^\circ$ from the Cu–O–Cu bond directions, while the observed inelastic stripes are rotated by $3^\circ$. The inelastic dispersion appears continuous and steep, consistent with earlier cuprate results [15,16]. However, the elastic signal shows a large and significant difference in $\delta_k$, appearing as a discontinuity in the dispersion relation at a vanishing energy transfer. Similar observations have been briefly remarked upon in underdoped La$_{2-x}$Sr$_x$CuO$_4$ ($x = 0.07$) [25] and La$_{2-x}$Ba$_x$CuO$_4$ ($x = 0.095$) [26]. In both cases, the observation was left unexplained.

To rule out that these surprising differences in $\delta_k$ and the stripe rotation are artifacts caused by experimental non-idealities, we have performed a virtual ray-tracing experiment in McStas [27,28]. The simulations use a precise model of the experiment including the guide system [29], monochromator, sample, analyzer, and detector; details can be found in Supplemental Material [10]. This method is known to accurately reproduce experimental effects like peak broadening and displacement [30]. The virtual experiments exclude misalignment of the instrument as a cause of the effect and show that the experimental resolution can cause a tiny shift in the observed incommensurability; see Fig. 3.

The experimentally observed shift in peak position is, however, more than an order of magnitude larger than what can be explained by instrument effects and is therefore a genuine property of the sample. Hence, in order not to violate the Goldstone theorem, the static and dynamic stripes must originate from different regions in the sample. There are two probable ways this can occur.

First, the dynamic stripes could be transverse fluctuations from the static stripe order, resembling ordinary spin waves. Because of the neutron scattering selection rules, the scattering observed in the elastic and inelastic channels stem from different twin domains as explained in detail in Supplemental Material [10]. This results in a shift in the observed peak position between the elastic and inelastic channels, comparable to the observed shift. The magnitude and direction of the shift due to twinning depend heavily on $\delta_h$ and $\delta_k$. Second, the static and dynamic spin response may originate from different microscopic regions which are not related by twinning. This suggests a real-space electronic phase separation of the crystal into regions with two different spin structures: one domain type which has static stripe order and associated dynamic stripes and another type of domain where only dynamic stripes are present.

At first glance, the twinning model seems to provide an explanation of our data. However, it fails to explain the similar observations in LBCO (where $\delta_h = \delta_k$) mentioned above [26], as the model requires $\delta_h < \delta_k$. Furthermore, the model relies on the assumption that the four twin domains display only one type of stripe order with associated transverse excitations. Most likely, these assumptions are too simplified, and relaxing any of them reduces the effect of twinning on the observed signal. We therefore turn to the second model: electronic phase separation.

Muon spin rotation experiments on highly oxygenated LCO + O show that the material electronically phase separates into a magnetic (A) and a superconducting (B) phase of roughly equal volume [31,32] and transition temperature $T_c \approx T_N \sim 40$ K with the present slow cooling conditions. Based on these experiments, we propose the following properties of the two phases.

Phase A is underdoped (resembling LSCO with $n_h = 0.125$) and has static magnetism (and weak fluctuations), responsible for the observed static signal and a small fraction of the dynamic signal. Phase B is optimally doped (resembling LSCO with $n_h = 0.16$) and superconducting with strong fluctuations, responsible for (the majority of) the observed dynamic signal.

We note that no spin gap was observed below $T_c$ in our experiments. An absence of a spin gap was also observed in the experiments on strongly underdoped LSCO [25] and LBCO [26], mentioned above. Both materials were suggested not to be $d$-wave superconductors but instead
Our results are consistent with this interpretation. A PDW state would require some degree of magnetic order in the SC phase, but this may be extremely weak and thus effectively invisible in our experiments. The simultaneous observation of gapless excitations and a shift in incommensurability in all three compounds suggests a connection between the two effects. The gapless excitations are likely a result of a PDW state in the sample, while the shift is caused by electronic phase separation. At present, it is unclear whether these two behaviors are related.

The critical temperature of the superconducting phase, \( T_c \), coincides with the Néel temperature of the magnetic phase, \( T_N \), such that above this temperature superconductivity and the static magnetism disappear, but strong stripe fluctuations remain. The fact that \( T_c \approx T_N \) is likely not coincidental, but it is unclear whether the electronic phase separation is caused by, or is the cause of, the close proximity of \( T_c \) and \( T_N \). We suggest a scenario where the ground state energies for phase \( A \) and phase \( B \) are very close and lowering the temperature below \( T_c \) will cause an electronic phase separation with concurrent static magnetism and superconductivity. The spatial distribution of impurity potentials as well as inhomogeneous hole doping become important parameters that can tip a region towards becoming type \( A \) or \( B \), e.g., by pinning.

The relative population of each phase is primarily controlled by the total number of holes but can also be influenced by an applied magnetic field or by crash cooling [11,34]. In the case of \( \text{LCO} + \text{O} \), crash cooling can further inflict a lowering of \( T_c \) which has been explained by a disconnection of the optimally superconducting pathways [35].

The crucial point is that, although the two phases are closely related, there is no \textit{a priori} reason why the stripe order in phase \( A \) and the stripe dynamics in phase \( B \) should have the same incommensurability. Indeed, our results show that this is not the case. This indicates that other properties of the stripes may not be identical either, and one should thus be extremely careful when interpreting neutron scattering data on stripes.

Phase separation has been suggested to occur in a number of cuprates or related compounds, a few of which we will mention here. \( \text{LSCO} \) with \( x = 0.12 \) has been suggested to phase separate into microscopic superconducting regions with gapped dynamic stripes and non-superconducting regions with static stripes [36]. Spontaneous, microscopic phase separation has also been observed in purely oxygen-doped \( \text{LCO} + \text{O} \) crystals [32] and in crystals doped with both oxygen and strontium [37]. Furthermore, recent studies of \( \text{La}_{\delta/3}\text{Sr}_{1/3}\text{CoO}_4 \) show evidence of microscopic phase separation into components with different local hole concentration [38,39]. In the latter material, the upper and lower parts of the hourglass dispersion are even proposed to originate from different nanoscale structures in the sample [38]. No discrepancy in the incommensurability between static and dynamic stripes was reported in these studies.

The idea of dynamic and static stripes having different origins is supported by a number of other observations. For example, the static and dynamic stripes exhibit different behaviors as a function of the temperature. In underdoped \( \text{LSCO} \) and in \( \text{LCO} + \text{O} \) as evidenced in this experiment, the static stripes vanish above \( T_c \), but the dynamic stripes remain to far higher temperatures [40–42]. In contrast, in the optimally doped region, the static stripes are altogether absent, while the dynamic stripes exist above a certain energy gap [16]. In the heavily overdoped region, it has been shown that substituting small amounts of Fe for Cu induces static magnetism [43]. The incommensurability of the induced magnetic order is governed by nesting of the underlying Fermi surface and differs from the 1/8 periodicity of the low-energy dynamic stripes.

When applying a magnetic field, the static stripes are, in general, strengthened [40,41,44–47], with a few exceptions [46,48]. In many cases, this happens with an accompanying change in the dynamic stripe spectrum [40,41,44], but in other cases, the dynamic stripe spectrum is unchanged [45]. Hence, the coupling between static and dynamic stripes is not simple and unique.

In conclusion, we have found that the dynamic stripes do not disperse towards the static stripes in the limit of vanishing energy transfer in a HTS compound. The effect is subtle and requires high flux and good resolution such as provided by the ThALES spectrometer in order to be observed. Our findings are, however, of prime importance, since they suggest that the observed static and dynamic stripes originate from different electronic phases in the sample, where one of these phases is likely to be a competitor for superconductivity with the development of static stripe order.

Our observations are relevant for all compounds displaying stripe order. As an example, the structurally similar, but nonsuperconducting compound \( \text{La}_{\delta/3}\text{Sr}_{1/3}\text{NiO}_4 \) (\( \text{LSNO} \)) displays magnetic and charge stripes with the dynamic stripes persisting at higher temperatures than the ordering temperature [49]. In some of these compounds, it has also been observed that the ordering vectors of the static and dynamic stripes do not coincide at vanishing energy transfer [50]. We speculate that a similar electronic phase separation could be at play here, as we suggest for \( \text{LCO} + \text{O} \). It is likely that this mechanism also explains earlier observations of unusual dispersions in \( \text{LSCO} \) [25] and \( \text{LBCO} \) [26]. Our findings may thus be a vital part in unveiling the nature of high-temperature superconductivity.

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