Structural Origin of Apparent Fermi Surface Pockets in Angle-Resolved Photoemission of Bi$_2$Sr$_{2-x}$La$_x$CuO$_{6+\delta}$

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Abstract

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Reference


DOI: 10.1103/PhysRevLett.106.127005
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We observe apparent hole pockets in the Fermi surfaces of single-layer Bi-based cuprate superconductors from angle-resolved photoemission. From detailed low-energy electron diffraction measurements and an analysis of the angle-resolved photoemission polarization dependence, we show that these pockets are not intrinsic but arise from multiple overlapping superstructure replicas of the main and shadow bands. We further demonstrate that the hole pockets reported recently from angle-resolved photoemission [Meng et al., Nature (London) 462, 335 (2009)] have a similar structural origin and are inconsistent with an intrinsic hole pocket associated with the electronic structure of a doped CuO$_2$ plane.

DOI: 10.1103/PhysRevLett.106.127005 PACS numbers: 74.72.Kf, 74.25.Jb, 74.72.Gh

The pseudogap is one of the defining properties of the hole-doped high-$T_c$ superconductors [1]. Understanding its origin is widely regarded as a key to unravelling the mechanisms of the high transition temperature superconductivity in these materials. While above the pseudogap temperature $T^*$ a large closed Fermi surface characteristic of ordinary metals is observed [2,3], at temperatures below $T^*$ angle-resolved photoemission (ARPES) measurements have long revealed disconnected Fermi “arcs,” centered around the $(0,0) \rightarrow (\pi, \pi)$ nodal direction [2,4–6]. Only recently, it has been claimed that distinct closed hole pockets can be observed by ARPES at particular compositions of the Bi$_2$Sr$_{2-x}$La$_x$CuO$_{6+\delta}$ (La-Bi$_2$201) cuprate [7]. While this would be qualitatively consistent with the demonstration of quantum oscillations in underdoped cuprates, indicative of small closed pockets in a high field [8–13], it remains unclear how it can be reconciled with numerous earlier ARPES studies.

A key difficulty is the structural complexity of Bi-based cuprates. They possess an orthorhombic lattice distortion [14], which is likely the origin of the so-called shadow Fermi surface [15–18]. In addition, they are prone to single [14,18,19] or even multiple [20] superstructure modulations along the crystallographic $b$ axis. These additional periodicities arise from a slight lattice mismatch between the BiO and CuO$_2$ planes and cause diffraction replica (DR) of electronic bands [21].

Here, we combine an extensive $k$-space survey of the Fermi-surface topology of La-substituted Bi2201 with a detailed structural analysis, in order to separate generic electronic effects from DR. We show that large superstructure periodicities of up to $14a_0$, coexisting with the already well-established periodicity of $\approx 4.2a_0$, are common. Such structural artifacts lead to imitations of closed Fermi-surface pockets in ARPES measurements. From their polarization dependence, we demonstrate that the front and back sides of the pockets derive from the main and shadow bands, respectively, and do not represent a single closed portion of Fermi surface intrinsic to the doped CuO$_2$ plane as claimed previously [7].

We investigated optimally (OP) and underdoped (UD) La-Bi$_2$201 with $x = 0.5$ (OP30K), $x = 0.75$ (UD20K), and $x = 0.8$ (UD14.5K) and optimally doped Bi$_{1.2}$Pb$_{0.35}$La$_{0.4}$Sr$_{1.4}$CuO$_{6+\delta}$ ([Pb-La]-Bi$_2$201, OP35K) samples. ARPES measurements on La-Bi$_2$201 (Figs. 1 and 3) were performed in the pseudogap phase at $\sim 17.5$ K and $\sim 33.5$ K for UD14.5K and OP30K La-Bi$_2$201, respectively, with linearly polarized He-I$_\alpha$ radiation ($\hbar\nu = 21.22$ eV) and a SPECS Phoibos 225 hemispherical analyzer, while the data on (Pb-La)-Bi$_2$201 (Fig. 2) were taken in the superconducting phase at 10 K with unpolarized light. The angular and energy resolutions for all measurements were set to $0.3^\circ$ and better than 20 meV, respectively. All low-energy electron diffraction (LEED) patterns shown here were recorded at temperatures between $\sim T_c$ and $\sim 2T_c$ with an incident electron energy of 35 eV.

Figure 1(a) shows the Fermi surface of UD14.5K La-Bi$_2$201 as measured by ARPES. Fermi arcs are visible centered around the nodal directions, although their intensity is suppressed along $\Gamma-Y$ due to matrix element effects which we shall discuss later. Multiple copies of these arcs...
can be observed, separated by $\sim 0.28 \, \text{Å}^{-1}$ along the $\Gamma - Y$ direction, consistent with DR corresponding to the established dominant superstructure periodicity of this material. In addition, weak features with opposite dispersion to the Fermi arcs appear to form several small closed pockets along the nodal direction (white arrows, $p_1 - p_3$). These features cannot be observed in optimally doped La-Bi$_2$2201, shown in Fig. 1(b), consistent with the findings of Meng et al. [7].

In the following, we show that the closed pockets appear as a natural consequence of structural complications in La-Bi$_2$2201. Lines of periodic diffraction maxima, characteristic of superstructure modulation along the $\Gamma - Y$ direction, are clearly discernible in the LEED pattern from OP30K La-Bi$_2$2201 [Fig. 1(d)]. From their spacing, the superstructure vector can be determined as $q_i = (q_1, q_2) \frac{a_i}{a}$ with $q_1 = 0.235 \pm 0.015$, in agreement with the DR in the ARPES and with earlier diffraction studies [22]. Intriguingly, LEED from UD14.5K La-Bi$_2$2201 [Fig. 1(c)] shows not only a similar superstructure vector $q_1 = 0.245 \pm 0.015$ but exhibits yet further diffraction maxima revealing the coexistence of a second superstructure with $q_2 = 0.130 \pm 0.015$. In order to demonstrate how these superstructure periodicities lead to the impression of hole pockets in ARPES, we first fit a tight-binding model to the Fermi surface of the main band for each doping [solid black lines in Figs. 1(a) and 1(b)] and then translate this band by $(\pi, \pi)$ to describe the shadow band resulting from the orthorhombic distortion. Finally, we add umklapp bands, that is, DR of the main and shadow bands, with the $q_i$ vectors determined independently from LEED. Without any further adjustment this simple model reproduces the entire measured Fermi surfaces for the UD14.5K and the OP30K samples over an extensive $k$-space range. In particular, it describes all apparent hole pockets in the underdoped sample and the absence of these pockets in optimally doped La-Bi$_2$2201. For example, the pockets $p_1$ and $p_2$ in UD14.5K La-Bi$_2$2201 are created by the $q_2$ DR of the shadow band crossing the main band, while another pocket ($p_3$) is formed by the $-q_2$ DR of the main band overlapping the $-q_2$ DR of the shadow band. All of these pockets are absent in OP30K La-Bi$_2$2201, which does not show the $q_2$ periodicity in LEED. It is therefore evident that the seemingly closed portions of Fermi surface in underdoped La-Bi$_2$2201 are not intrinsic but appear from the overlapping of DR resulting from multiple superstructure periodicities.

We stress that the appearance of these pockets is not directly tied to the hole concentration, or the presence of a pseudogap, but rather to the particular structural modulations. Indeed, we also observe multiple superstructures, with vectors of $q_1 = 0.225 \pm 0.015$ and $q_2 = 0.072 \pm 0.015$, in optimally doped (Pb,La)-Bi$_2$2201 [Fig. 2(a)]. Although Pb doping tends to suppress superstructure-related features in ARPES from Bi-based cuprates, the measured Fermi surface [Fig. 2(b)] clearly shows the presence of umklapp resulting from these superstructure vectors and their combinations, consistent with a tight-binding model using the $q_i$ vectors determined from LEED. Similar

![Image](https://example.com/figure1.png)

FIG. 1 (color online). ARPES Fermi surface of (a) UD14.5K and (b) OP30K La-Bi$_2$2201, measured with $p$ polarization. Extracted contours (blue circles) and a tight-binding model of the main (solid line), shadow (dashed line), and $\pm q_1$ or $\pm 2q_1$ (green), $\pm q_2$ (blue), and $\pm (q_1 + q_2)$ (red) umklapp bands are overlaid on the data. LEED from (c) UD14.5K and (d) OP30K La-Bi$_2$2201. A magnified view of the blue region and a line cut along the $\Gamma - Y$ direction (red line) are shown to the left of and below each pattern, respectively. Extracted peak positions (red dots) and those expected for superstructure peaks at $k_0 \pm m q_i \pm m_2 q_2$ (vertical gray lines) are in good agreement. The inset in (a) shows a simplified Fermi surface with only the main (MB) and shadow (SB) bands.
In contrast, the shadow band, which has the opposite parity of the main band [15], is visible in \( p \) polarization but not in \( s \) polarization. This switching of intensities is clearly seen for OP30K La-Bi2201 in Fig. 3(a) [23]. For measurements in the 2nd Brillouin zone, along the cut shown in Fig. 3(d), the polarization is no longer strictly \( s \) or \( p \) since the sample is tilted off-vertical by \( \sim 30^\circ \). Nevertheless, a strong relative intensity variation can still be observed between the main and shadow bands on switching from dominant \( p \) polarization \((I_{MB} > I_{SB})\) smaller) to \( s \) polarization \((I_{MB} < I_{SB})\) larger), as shown in Fig. 3(b). The equivalent dispersion measured in UD14.5K La-Bi2201 is shown in Fig. 3(c). With \( p \) polarization, two strong dispersions can be seen due to the main band and its \( -\mathbf{q}_1 \) DR, with two weaker neighboring bands which form the back side of the apparent Fermi-surface pockets (marked in the Fermi level momentum distribution curve by circles and crosses, respectively). This gives the appearance that the pocket on the Fermi surface is holelike, as claimed in Ref. [7]. However, on switching to \( s \) polarization, these “pocket-forming” bands are strongly suppressed relative to the main bands, as is the spectral intensity of the back side of all of the pocket features which can be seen in the ARPES Fermi surface [Fig. 3(d)]. This switching of intensities due to different parities of the front and back sides of the Fermi-surface pockets is difficult to reconcile with intrinsic pockets of a reconstructed Fermi surface and confirms that these features are derived from the shadow band.

Given this, one must revise the conclusions of Ref. [7] regarding intrinsic hole pockets. Meng et al. [7] considered only umklapp bands arising from a \( q_1 = 0.24 \) superstructure modulation and found that these DR could not explain their data. However, LEED from La-Bi2201 with very similar composition to the UD18K sample of Ref. [7] shows not only the \( q_1 = 0.24 \) superstructure but also a second superstructure with \( q_2 = 0.12 \pm 0.015 \) [see

FIG. 2 (color online). (a) LEED pattern and (b) ARPES Fermi surface of OP35K (Pb,La)-Bi2201, showing the presence of two superstructure vectors, and the resulting apparent pockets in the Fermi surface. A tight-binding model, using superstructure vectors determined from the LEED analysis, is also shown in (b).

to the situation in underdoped La-Bi2201, the overlapping of several of these bands gives rise to features that appear as closed Fermi-surface pockets. However, as for UD14.5K La-Bi2201, this is entirely due to structural effects and should not be confused with either an intrinsic hole pocket or an incommensurate density-wave order.

The structural origin of the observed pockets can be further confirmed by polarization-dependent ARPES measurements, as shown in Fig. 3. The \( x^2 - y^2 \) symmetry of the hole in the Cu \( d \) shell is odd with respect to the \( \Gamma - Y \) direction. Consequently, for the experimental geometry employed here, with the incident light and detected electrons both within the horizontal plane, the main band can be observed when measuring with \( s \)-polarized light along the \( \Gamma - Y \) azimuth but is suppressed for \( p \) polarization. This is the reason why the intensity along the Fermi arc diminishes approaching the \( \Gamma - Y \) nodal line in Figs. 1(a) and 1(b). In contrast, the shadow band, which has the opposite parity

FIG. 3 (color online). Polarization dependence of ARPES. (a) Dispersion along \( \Gamma - Y \) of OP30K La-Bi2201. (b),(c) Dispersion close to the equivalent direction in the 2nd Brillouin zone [along dashed lines in (d)] of OP30K and UD14.5K La-Bi2201. (d) ARPES Fermi surface of UD14.5K La-Bi2201. All spectra were measured with \( p \) (top) and \( s \) (bottom) polarization.
We note that our findings do not exclude the presence of intrinsic Fermi-surface pockets in cuprates, where the back side of the pocket has negligible spectral weight in ARPES measurements [26]. However, we remark that, to date, clearly discernible hole pockets have been reported only from ARPES measurements in Bi-based and La-based cuprate systems [7,27], which are both subject to structural distortions. No such observation has been made in compounds free of such distortions, for example, CCOC [5] and YBCO [28].

We acknowledge financial support from the Scottish Funding Council, the United Kingdom EPSRC (Grant No. EP/F006640), the ERC [St Andrews], the Killam program, the Sloan Foundation, the CRC program, NSERC, CFI, CIFAR Quantum Materials, BCSI [UBC], TRF-OHEC [Suranaree], and NSF Grant No. DMR-0847345 [Cornell].

Fig. 4(a)]. In Fig. 4(b), we show that the measured pockets, and indeed all features of the Fermi-surface topology extracted in Ref. [7], are accurately described by a tight-binding model of only the conventional main and shadow bands, provided DR are included corresponding to both of these superstructure vectors. Together with the polarization-dependent ARPES presented above, this shows unambiguously that the observations of Meng et al. [7] have a trivial interpretation and cannot be taken as evidence for elusive Fermi pockets.

We also note that a structural origin of the pockets explains several puzzling observations of Ref. [7]. First, the main band (Fermi arc) was observed to be longer than the back side of the claimed pocket [as is also evident here in Figs. 1(a) and 2(b)]. The lower intensity of the shadow band and its DRs, which appear to “close” the pockets, compared to the main band naturally accounts for this seemingly contradictory coexistence of Fermi arcs and hole pockets. Second, the spectral weight of the back side of the pockets in Ref. [7] appears largest close to the nodal line. This is in contrast to theoretical expectations for an intrinsic pocket [24,25] but consistent with a superstructure replica of the shadow band. Third, multiple hole pockets were observed in Ref. [7], attributed to a q = 0.24 superstructure replica of a single intrinsic pocket. However, the front- to back-side spectral weight ratio differs for these pockets. Again this suggests that the two sides of the pockets derive from different bands, namely, the main and shadow band, whose umklapps display complex intensity variation due to matrix element effects. In addition, the superstructure vectors depend sensitively on doping as shown from the LEED analysis presented here, which provides a simple explanation for the unusual doping dependence of the hole pockets reported by Meng et al. [7].

We acknowledge financial support from the Scottish Funding Council, the United Kingdom EPSRC (Grant No. EP/F006640), the ERC [St Andrews], the Killam program, the Sloan Foundation, the CRC program, NSERC, CFI, CIFAR Quantum Materials, BCSI [UBC], TRF-OHEC [Suranaree], and NSF Grant No. DMR-0847345 [Cornell].

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21. Throughout, we distinguish the (, ) umklapp of the main band, giving rise to the shadow band, from the superstructure diffraction replica to be consistent with conventional terminology.
23. The lack of complete suppression of the forbidden transitions is due to the finite degree of polarization (~80%) of the incident light in the experimental setup used here.