Three Dimensionality and Orbital Characters of the Fermi Surface in (Tl,Rb)$_y$Fe$_{2-x}$Se$_2$


Phys. Rev. Lett. 109, 037003 — Published 20 July 2012

DOI: 10.1103/PhysRevLett.109.037003
Three-dimensionality and orbital characters of Fermi surface in (Tl,Rb)$_2$Fe$_{2-x}$Se$_2$


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We report a comprehensive angle-resolved photoemission spectroscopy (ARPES) study of the tridimensional electronic bands in the recently discovered Fe-selenide superconductor (Tl,Rb)$_2$Fe$_{2-x}$Se$_2$ ($T_c = 32$ K). We determined the orbital characters and the $k_z$ dependence of the low-energy electronic structure by tuning the polarization and the energy of the incident photons. We observed a small 3D electron Fermi surface pocket near the Brillouin zone (BZ) center and a 2D like electron Fermi surface pocket near the zone boundary. The photon energy dependence, the polarization analysis and the LDA calculations suggest a significant contribution from the Se 4$p_x$ and Fe 3$d_{xy}$ orbitals to the small electron pocket. We argue that the emergence of Se 4$p_x$ states might be the cause of the different magnetic properties between Fe-chalcogenides and Fe-pnictides.

PACS numbers: 74.70.Xa, 74.25.Jb, 79.60.-i, 71.20.-b

The recently discovered Fe-chalcogenide superconductors $A_y$Fe$_{2-x}$Se$_2$ ($A$=K, Rb, Cs, Ti etc.) with maximum $T_c = 32$ K [1, 2] changed the landscape of Fe-based superconductivity. Unlike the other Fe-based superconductors, for which both hole-like and electron-like Fermi surfaces (FSs) are observed simultaneously [3–9], these new materials are characterized by the absence of hole-like FS [10–13]. As with most of the other materials though, nearly isotropic superconducting (SC) gaps in the strong coupling regime are found on the observed FSs [11–13], suggesting a possible universal pairing mechanism for all Fe-based superconductors. Local antiferromagnetic (AFM) exchange interactions emerge as a serious candidate for the unification and the $A_y$Fe$_{2-x}$Se$_2$ system constitutes a good test to this approach. Indeed, neutron scattering experiments [14, 15] suggested a magnetic ground state of the parent compound that differs from the collinear order commonly observed in the Fe-pnictides [16, 17]. In addition, large AFM exchange coupling constants have been determined experimentally in $A_y$Fe$_{2-x}$Se$_2$ as well as in FeTe [15, 18] for the next ($J_2$) and next-next ($J_3$) nearest neighbors. Unfortunately, there is no experimental understanding of the origin of these enhanced constants that tune the magnetic ground state of these materials and may as well tune their SC properties.

High quality single crystals with nominal composition (Tl,Rb)$_{0.8}$Fe$_{1.63}$Se$_2$ were synthesized by the flux method. The SC transition temperature $T_c = 32$ K with a transition width of 2 K was measured with a PPMS [2]. Samples were mounted in an inert gas protected glove box to prevent reaction with moisture. Samples with size smaller than $1 \times 1 \text{ mm}^2$ were cleaved in situ, yielding a flat mirror like (001) surface. ARPES measurements were performed at Renmin University of China with a Scienta R3000 analyzer and at the Apple-PGM and PGM beamlines of the Synchrotron Radiation Center (Stoughton, WI), with Scienta 200U and Scienta R4000 analyzers, respectively. Normal state spectra were taken at $T = 40$ K. During measurements, the pressure was maintained below $4 \times 10^{-11}$ Torr and no noticeable aging was observed during measurement cycles. The $E_F$ was referenced to a fresh Au polycrystalline film in good electrical contact to the sample.

The unfolded BZ of (Tl,Rb)$_2$Fe$_{2-x}$Se$_2$ and high symmetry points are presented in Fig. 1(a) and the experimental setup is shown in Fig. 1(d). The band dispersion $E_B(k)$ along the $c$-axis can be measured by changing the photon energy ($h\nu$) with $k_z$ estimated using the formula [19, 20]:

$$k_z = \frac{1}{\hbar} \sqrt{2m \nu [(h\nu - \phi - E_B + V_0) - \hbar^2 k^2_y]} ,$$

where $\phi$ is the work function and $V_0$ the inner potential of the sample. In (Tl,Rb)$_2$Fe$_{2-x}$Se$_2$, with the lattice constant $c = 14.56$ Å [21], $\phi = 4.3$ eV and empirically determined $V_0 = 15$ eV, we found that $h\nu = 18$, 38 and 63 eV are close to the $\Gamma$ point, while 28, 50 and 78 eV are close to the $Z$ point.

Figs. 1(b) and 1(c) compare the normal state FSs obtained in the $\Gamma$ plane ($h\nu = 38$ eV) and in the $Z$ plane ($h\nu = 28$ eV), respectively. The FSs are in agreement with previous reports on the electronic structure of $A_y$Fe$_{2-x}$Se$_2$ [10–13]. We identify a FS near the BZ boundary (M, A), which is almost $k_z$-independent.
FIG. 1. (Color online) (a) Schematic unfolded BZ of (Tl,Rb)$_3$Fe$_{2−x}$Se$_2$. (b)-(c) Integrated intensity plots at $E_F$ [-10 meV, 10 meV] of (Tl,Rb)$_3$Fe$_{2−x}$Se$_2$ in the Z ($h\nu=28$ eV) and $\Gamma$ ($h\nu=38$ eV) planes, respectively. Dashed lines are extracted FSs. (d) Sketch of the experimental polarization setup. The $s$ and $p$ polarizations are for electric fields perpendicular and parallel to the mirror plane (cyan color), respectively. We note that our experimental setup for the $s$ geometry includes a polarization component along $z$. (e) Second derivative $[\partial^2 I(\omega, \kappa)/\partial \omega^2]$ plots of band dispersion along $\Gamma$–M, indicated as cut 1 in (b). (f) Corresponding EDCs. Dashed lines are guides to the eye for the band dispersions. (g) and (h) Same as (e) and (f) but along Z–A.

This observation is supported by the band dispersions, which are given in Figs. 1(c)-(f) and 1(g)-(h) for the $\Gamma$–M and Z–A high symmetry lines, respectively [24]. Compared with the Fe-pnictides, the hole-like bands near the 2D zone center are shifted down to about 100 meV below $E_F$ and thus do not contribute to the FS, as reported previously [10–13].

The main contrast between the $\Gamma$ and Z plane lies in the existence of an additional Z-centered electron-like band, $\kappa$, which is observed in the Z plane and fully absent in the $\Gamma$ plane. In the Z plane, the $\kappa$ band has its minimum at 20-30 meV below $E_F$ and crosses $E_F$ at $k_\parallel \approx 0.1\pi/a$, yielding the small Z-centered FS pocket observed in Fig. 1(c). To investigate the detailed band dispersion along the $k_z$ direction, we did ARPES measurements with $h\nu$ ranging from 15 to 80 eV, which covers $k_z$ from $Z_1$ to $Z_3$. Fig. 2 shows the $h\nu$–$k_\parallel$ FS intensity plot, with $k_\parallel$ oriented along the Z–A line. We observe three FSs centered at 28 eV ($Z_1$), 50 eV ($Z_2$) and 78 eV ($Z_3$) respectively, with the FSs lying halfway between Z and $\Gamma$. We note that the total enclosed FS area of the $\kappa$ and $\delta$ bands is about 6.5% of the unfolded BZ (13% electron per Fe), confirming electron-doping in this system.

The $h\nu$ variation of the $\kappa$ band is also well illustrated by the intensity plot of EDC curvature [25] along (0, 0, $k_z$), as shown in Fig. 2(b). The $\kappa$ band has a clear dispersion along the $k_z$ ($h\nu$) direction, with its band minimum at Z ($Z_1$ and $Z_2$) and across $E_F$ between Z and $\Gamma$. Interestingly, a direct comparison of the EDCs at 28 eV ($Z_1$), 50 eV ($Z_2$) and 78 eV ($Z_3$), displayed in Fig. 2(c), reveals that the spectral intensity associated with the $\kappa$ band decreases with $h\nu$. Besides the $\kappa$ band, the $\omega$ band lying around 350–450 meV below $E_F$ also exhibits appreciable $h\nu$ variation, with a band maximum at Z and a minimum at $\Gamma$. However, its spectral intensity evolves just the opposite way, with a stronger peak at $Z_3$ than at $Z_2$ and $Z_1$. This has an important consequence: although the strongly $k_z$-dispersive nature of these bands indicate that they are unlikely related to a surface state and that they carry significant components from orbitals extended along the c-axis, their respective dominant orbital or elemental characters must be different. The photoemission cross sections [26] of the Fe 3d and Se 4p states within the $h\nu=20–80$ eV range are plotted in Fig. 2(d). In contrast to the spectral weight of the $\omega$ band, the $h\nu$ variation of the spectral weight of the $\kappa$ band, as shown in the inset of Fig. 2(d), cannot be described by the cross section calculated for Fe 3d states. Surprisingly, it shows much better agreement with the cross section calculated for Se 4p states, which monotonically decreases in the considered $h\nu$ range.

To refine our assumptions on the nature of the electronic states in the vicinity of $E_F$, we performed a photon
polarization dependence study. Fig. 3 shows the bands in the Z plane with s and p polarized incident photons, where s and p respectively refer to an electric field perpendicular and parallel to the photoemission plane defined by the normal of the sample surface and the photoemitted electrons. (we note that our s experimental geometry contains a component of the light polarization along the z-axis). Along the Z(α–A(β) direction, the α band is visible using p polarization but suppressed with s polarization. The β band behaves oppositely. By analogy with the Ba0.4K0.6Fe2As2 and FeTe0.55Se0.45 compounds [27], this suggest that the the α and β bands are likely related to the Fe 3dxy and the even combination of the Fe 3dxz and Fe 3dyz orbitals, respectively. The electron-like band δ at the zone boundary A, with a minimum at ~50 meV, is seen in both configurations, suggesting a complicated orbital character that deserves further detailed study. While the κ band at Z1 is clearly observed with s polarized light, as shown in Figs. 3(a) and 3(b), it is barely detectable with pure p polarization [see Figs. 3(c) and 3(d)]. The photon polarization dependence of the κ band is consistent with either a wave function having odd symmetry with respect to the measurement plane or with a z-oriented wave function.

To clarify the situation, we present in Fig. 4 our LDA calculations for TlFe2Se2 using the experimental structure parameters [28], along with projections for the Se 4pz, Fe 3dxy and Fe 3dxz,3dyz orbitals. As with previous LDA calculations [29–32], we notice discrepancies with ARPES observations [10–13]. The main one is a downshift of the hole bands near the zone center accompanied by an upshift of the electron bands near the zone boundary, which has been explained in terms of correlation effects and the evidence of interband coupling [10, 33]. Nevertheless, LDA captures several important features of the ARPES observations [10]. LDA calculations indicate that the energy of the κ band is sensitive to the height of the Se atoms and shifts down below E_F in (Tl,Rb)Fe2−xSe2 [29]. Interestingly, the LDA orbital projections suggest that in addition to the Se 4pz orbital character, the κ band also carries a non-negligible Fe 3dxy component, thus explaining its sensitivity to odd
configurations of light polarization. The calculations also suggest that the \( k_z \) dispersive \( \omega \) band at higher binding energy origins mainly from the Fe 3d_{x^2−y^2} orbital.

It has been recently proposed that local AFM exchange fluctuations [34], which are characterized by exchange coupling constants, mainly determine the momentum distribution of the SC gap in high-temperature superconductors such as cuprates and Fe-based superconductors [35]. The \( \text{A}_{x}\text{Fe}_{2−x}\text{Se}_2 \) system has a complicated magnetic structure that differs from that of the Fe-pnictides [14, 15, 36]. A recent neutron scattering study reported a ferromagnetic nearest neighbor exchange coupling \( (J_1) \), appreciable AFM next-nearest \( (J_2) \) and next-next-nearest neighbor couplings \( (J_3) \), as well as an AFM interlayer coupling \( (J_p) \) in \( \text{Rb}_n\text{Fe}_{2−x}\text{Se}_2 \) [15]. It is widely believed that the magnetic super-exchange interactions in Fe-based superconductors are at least partially As/Se-bridged [37]. Our experimental observation of \( \text{Se} 4p_z \) states hybridized with Fe 3d orbitals at \( E_F \) in \( \text{(Ti,Rb)}_n\text{Fe}_{2−x}\text{Se}_2 \) provides a natural explanation to the existence of a large \( J_3 \) AFM exchange coupling. An intuitive picture can be argued as follows. Let’s consider a Fe atom. A local Wannier state can be created because the Fe 3d_{x^2y^2} orbital strongly hybridizes with the 4p_z orbital of its four Se atom neighbors, as illustrated in Fig. 4(d). This Wannier state can link the two neighboring Fe atoms on each side through a standard super-exchange interaction, thus resulting in the \( J_3 \) AFM coupling. In this picture, the \( J_3 \) is very sensitive to the hybridization due to the size of the \( p_z \) orbital and the distance of the Se/Te atoms from the Fe-layers. Interestingly, a possible hybridization between Fe 3d and \( p_z \) states has also been noted in the cousin \( \text{FeSe}_{0.45}\text{Te}_{0.55} \) system [6], which is also characterized by a large \( J_3 \) AFM coupling.

To summarize, we performed ARPES measurements on SC \( \text{(Ti,Rb)}_n\text{Fe}_{2−x}\text{Se}_2 \) with various \( h \nu \) values and different photon polarizations. We observed a small \( Z \)-centered electron-like FS pocket. LDA calculations and experimental data suggest that it originates mainly from a mixture of the \( \text{Se} 4p_z \) and Fe 3d_{x^2y^2} orbitals. These orbitals at \( E_F \) enhance the exchange coupling between Fe ions bridged by Se and may play an important role in the superconductivity of the \( \text{A}_x\text{Fe}_{2−x}\text{Se}_2 \) family.

This work was supported by the grants from the Chinese Academy of Sciences (2010Y1JBJ6), the Ministry of Science and Technology of China (2010CB923000, 2012CB821400 and 2011CBA001001), the National Science Foundation of China (10774189, 10974175, 11004232, 11190024 and 11050110422) and the Research Funds of Renmin University of China (12XNH093). This work is based in part upon research conducted at the Synchrotron Radiation Center which is primarily funded by the University of Wisconsin-Madison with supplemental support from facility Users and the University of Wisconsin-Milwaukee.

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[24] We note that we did not observe the FS reconstruction due to the \( \sqrt{5} \times \sqrt{5} \) magnetic and structural reconstruction reported by neutron scattering.