Temperature-Dependent Fermi Surface Evolution in Heavy Fermion CeIrIn$_5$

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We address theoretically the evolution of the heavy fermion Fermi surface (FS) as a function of temperature ($T$), using a first principles dynamical mean-field theory approach combined with density functional theory. We focus on the archetypical heavy electrons in CeIrIn$_5$. Upon cooling, both the quantum oscillation frequencies and cyclotron masses show logarithmic scaling behavior [$\sim \ln(T_0/T)$] with different characteristic temperatures $T_0 = 130$ and 50 K, respectively. The enlargement of the electron FSs at low $T$ is accompanied by topological changes around $T = 10$–50 K. The resistivity coherence peak observed at $T \approx 50$ K is the result of the competition between the binding of incoherent $4f$ electrons to the sp$d$ conduction electrons at Fermi level ($E_F$) and the formation of coherent $4f$ electrons.

In cerium-based heavy electron materials, the $4f$ electron's magnetic moments bind to the itinerant quasiparticles to form composite heavy quasiparticles at low temperature ($T$). The volume enclosed by the Fermi surface (FS) in the Brillouin zone incorporates the moments to produce a "large FS" due to the Luttinger theorem. When the $f$ electrons are localized free moments, a "small FS" is induced since it contains only broad bands of conduction sp$d$ electrons. So, the FS volume is a sensitive probe of the character, localized or itinerant, of the heavy fermion system.[1]

Intensive efforts have been devoted to the study of the quantum phase transition leading from a small to large FS at strictly zero temperature. While the FS, as a surface of discontinuity in the momentum distribution function, is sharply defined only at zero temperature, experimental probes such as the angle-resolved photoemission spectra (ARPES) and magnetic quantum oscillation experiments such as de Haas–van Alphen (dHvA) or Shubnikov–de Haas experiments identify the region of momentum space where zero energy fermionic excitations exist at finite temperature. ARPES experiment directly observes the FS in the momentum space. But high resolution is required to determine the FS size. The quantum oscillation experiments measure the precise value of the FS area in a specific plane by probing the oscillation frequencies of magnetization as a function of the applied magnetic field. The quantum oscillation frequency ($f$), the so-called dHvA frequency, is proportional to the extremal cross-sectional area $S_F$ of the FS ($F = \hbar S_F/2\pi e$). The quantum oscillation experiments also provide information on the cyclotron effective electron mass $m^* = (\hbar^2/2\pi)\delta S_F/\delta\omega$ and the geometry of the FSs.

The band structure calculation is a complementary tool to the quantum oscillation experiment to analyze the complicated FS of the multiple band system. Quantum oscillation frequencies of heavy fermion materials, such as CeCu$_6$, UPt$_3$, and Ce(Fe,Co)In$_5$, are explained well by conventional band calculations because the itinerant $4f$ electrons behave as conduction electrons near $E_F$. Although the geometry and volume of FSs are well explained by the density functional theory (DFT) band calculation, the detected cyclotron mass $m^*$ is much larger than the corresponding DFT band mass $m_b$.[2–5], because the DFT calculation cannot describe the correlated $4f$ electronic states correctly. When the $4f$ electrons are localized in the antiferromagnetic (AFM) compounds, such as CeRhIn$_5$, CeIn$_3$, CeRh$_2$Si$_2$, the $4f$-localized band model is more applicable to the description of the quantum oscillation experiments.[2,5]. The $4f$-localized band model can be performed by treating the $4f$ electrons as core within the DFT (open-core DFT) band calculation[2] or by employing the DFT + $U$ band method ($U$ is the on-site Coulomb interaction)[6].

CeTIn$_5$ ($T = \text{Co, Rh, and Ir}$) has been a prototypical system to study the crossover behavior between the itinerant and localized $4f$ electrons. CeCoIn$_5$[7] and CeIrIn$_5$[8] have itinerant $4f$ electrons and superconducting ground states at low $T$. On the other hand, CeRhIn$_5$ has localized $4f$ electrons and the AFM ground state. The measured dHvA frequency of each compound identifies the nature of Ce $4f$ electrons, whether they are itinerant or localized. CeCoIn$_5$[2,4,9] and CeIrIn$_5$[2,3] have enlarged electron FSs due to the contribution of itinerant $4f$ electrons, while CeRhIn$_5$ has similar geometry of FSs but smaller size of FSs[2,5]. For CeRhIn$_5$, pressure-induced superconductivity was observed for $P > 1.63$ GPa[10], and the drastic change in the FS was detected at a critical pressure of $P_c = 2.35$ GPa[11]. On the other hand, CeRh$_{1-x}$Co$_x$In$_5$ shows the doping-dependent reconstruction of FS deep inside the magnetically ordered state[12], away from the quantum critical point (QCP). The single-site dynamical mean-field theory (DMFT) is known to be a proper approach for the
description of finite temperature properties near the QCP. According to recent DMFT approaches [13], the single-site DMFT study gives a correct result well above a Neél order due to the suppression of the nonlocal correlation. The intersite correlation becomes crucial only at extremely low $T$ near the Neél ordering. Because CeIrIn$_5$ does not show the AFM ground state and the temperature in our calculation is usually well above the nonlocal fluctuation region, our DMFT study is still valid for the description of the system near the QCP. The $T$-dependent evolution between itinerant and localized electrons has also been described by the phenomenological two-fluid model, where the universal scaling behavior can be applied to various physical properties of the heavy fermion compounds [14–16].

In this Letter, we have addressed the $T$-dependent crossover from localized to itinerant $4f$ electrons in CeIrIn$_5$, and investigated its effects on the FS properties and electrical resistivity. The charge self-consistent version of the DFT + DMFT approach [17], as implemented in Ref. [18], is based on the full-potential linearized augmented plane-wave band method [19]. The correlated $4f$ electrons are treated dynamically by the DMFT local self-energy, while all other delocalized $spd$ electrons are treated on the DFT level. The local self-energy matrix $\Sigma(\omega)$ is calculated from the corresponding impurity problem, in which full atomic interaction matrix is taken into account [20]. To solve the impurity problem, we used the vertex corrected one-crossing approximation (OCA) [17] and confirmed that the low $T$ limit of the OCA is consistent with the continuous time quantum Monte Carlo method [21,22]. (More information of the computational algorithm is given in the Supplemental Materials [23]).

The main difference between low and high $T$ spectral functions in the DFT + DMFT calculation is the existence of $4f$ bands near $E_F$, as shown in Fig. 2 of the Supplemental Materials [23]. Ce $4f$ bands at high $T$ are absent near $E_F$, and their spectral weights are distributed into the lower and upper Hubbard bands. The spectral function near $E_F$ can be well described by the quasiparticle band structures of other $spd$ electrons although there is a small scattering rate due to the hybridization between the conduction electrons at $E_F$ and the incoherent Ce $4f$ electrons in the Hubbard bands. As decreasing $T$, the spectral weight of the renormalized Ce $4f$ bands is increased continuously (see Movie 1 in the Supplemental Materials [23]). The hybridization of the $4f$ and other $spd$ bands produces very massive almost flat quasiparticle band structures near $E_F$. These flat bands emerge as the narrow Kondo peak at $E_F$ in the photoemission spectra [24].

The $T$-dependent FS has been extracted from the quasiparticle band structures. At low $T$, the FSs of the DFT + DMFT calculation are very similar to those of the DFT calculation, as shown in Fig. 1. Upon heating, the Ce $4f$ electrons become localized and their contribution to $E_F$ is suppressed. Accordingly, the areas of electron FSs ($\alpha_i$ and $\beta_i$) are continuously decreased. In contrast, there occur rather big changes in other FS areas upon heating. The areas of the $g$ and $h$ hole FSs on the $z = 0$ plane grow and merge into one closed electron FS. The $a$ electron FS identified at $T = 10$ K on the $z = \pi$ plane is divided at high $T$, and so new hole FSs appear near $Z$ and $R$ symmetry points. The continuous $T$-dependent evolution of the FS is provided in Movie 2 in the Supplemental Materials [23]. By integrating the volume of electron FSs, the occupancy of the conduction electrons has been counted. It shows the continuous change from 3 to 4 electrons as temperature is decreased, which reflects the participation of one Ce $4f$ electron in the bonding.

Because the area of the FS is directly related to the size of the $4f$ electron contribution to $E_F$, we have investigated the $T$-dependent dHvA frequencies, as shown in Fig. 2(a). At high $T$, the dHvA frequencies are well consistent with those from the Ce $4f$ open-core DFT calculation. With decreasing $T$, they show the continuous increase with the participation of $4f$ electrons to $E_F$ and follow the scaling behavior of $\ln(T_0/T)$, as shown in Fig. 2(c). All the branches show the same characteristic
The number of conduction electrons increases with decreasing temperature. The cyclotron mass corresponds to the effective mass of carriers at the specific FS. As shown in Figs. 2(b) and 2(d), the calculated cyclotron masses also increase upon cooling and follow a similar scaling behavior of \( \ln(T_0/T) \) with \( T_0^m \sim 50 \) K. The calculated cyclotron masses are also well fitted with the two-fluid model by Yang and Pines [15]: 
\[
(1 - T/T_0)^{3/2}[1 + \ln(T_0/T)]
\] with the same \( T_0^m \sim 50 \) K, as shown in Fig. 3 in the Supplemental Materials [23]. Interestingly, \( T_0^m \) is coincident with the coherent temperature \( T^* \) of Ce 4f states [24], but clearly different from \( T_0^f \). This feature reveals that the 4f electrons start to participate in bonding through the hybridization with \( spd \) electrons at the temperature scale \( T_0^f \), which is higher than the temperature \( T_0^m \) at which the coherent heavy fermion electronic states are formed. These results are reminiscent of recent experiment, which shows the occurrence of FS reconstruction much earlier than the quantum critical transition [12]. Note that the above scaling law is consistent with the two-fluid model [15], in which the coherent 4f bands start to grow below \( T^* \). All the calculated cyclotron masses at \( T = 10 \) K seem to be overestimated with respect to the experimental values [3] roughly by a factor of 2. It is well known that the value

FIG. 2 (color online). The \( T \)-dependent dHvA frequencies (F) and cyclotron effective masses (\( m^* \)). The dHvA frequencies (a) and effective masses (b) of \( \alpha_i \) and \( \beta_i \) branches are obtained from the DFT + DMFT method and compared with those from the DFT method and experiments (exp.). The corresponding FSs for each branch are provided in Figs. 1(a) and 1(e), except \( \alpha_i \) that corresponds to the maximum frequency among \( \alpha_i \) branches and is located between the \( z = 0 \) and \( z = \pi \) planes. At high \( T \), the cyclotron masses are very small, ranging from 0.4 to 0.7\( m_0 \) (\( m_0 \) is the bare electron mass) for \( \alpha_i \) and \( \beta_i \) branches. Such small cyclotron masses are also reproduced in the 4f open-core DFT calculation, in which only dispersive \( spd \) bands are crossing \( E_F \). The low \( T \) dHvA frequencies from the DFT + DMFT method are consistent with the results of DFT method in which the 4f electrons are considered as itinerant type. (c) The renormalized \( \Delta F \) of each branch shows the scaling behavior of \( \ln(T_0/T) \) with the characteristic \( T \) of \( T_0^f \sim 130 \) K. (d) All the renormalized \( \Delta m^* \)'s show the similar scaling behavior, but with \( T_0^m \sim 50 \) K.
of cyclotron mass has a substantial dependence on the applied magnetic field [4]. It is noted that the suppression of the $m^*$ with increasing magnetic field was studied by the spin-dependent mass [25–28] or the Zeeman effect [29,30].

The continuous change of FS properties with $T$ variation is deeply related to the transport properties. Figure 3 provides the calculated resistivity for CeIrIn₅ as a function of $T$, which is compared to the experimental electrical resistivity. The electrical resistivity is calculated using the real part of the dc conductivity ($\sigma$) [18] based on the DFT + DMFT spectral function near $E_F$:

$$\sigma^{\mu\nu} = \frac{e^2}{V} \sum_k \int d\omega \left( -\frac{df}{d\omega} \right) \text{Tr}[A(k, \omega)\nu^{k\mu}A(k, \omega)\nu^{k\nu}].$$

Here $\mu$ and $\nu$ represent spatial coordinates. $V$, $f(\omega)$, and $\nu$ are the primitive volume, the Fermi Dirac distribution function, and the velocity, respectively. The calculated resistivities from low to high $T$ are in good agreement with the experimental resistivity. At high $T$, the electronic carriers from dispersive $spd$ bands become more and more decoupled from localized electrons in the $4f$ shell; hence, the carriers are scattered less at very high $T$. Upon cooling, the hybridization among local moments and $spd$ carriers increases while the $4f$ electrons remain very incoherent above 50 K, causing enhanced scattering mechanism for electric carriers. Below the scale $T_{m}^0$, the electrons in the $4f$ shell also gain coherence which substantially suppresses the resistivity. Therefore, the maximum resistivity is observed near 50 K. Insets (a), (b), and (c) of Fig. 3 show the broadening of spectral weight at $E_F$, calculated at low, crossover, and high $T$, respectively. The broadening corresponds to the scattering rate at the specific $k$ point. It is noted that the spreading of the spectral weight at crossover is wider than that at high or low $T$. This finding confirms that the DFT + DMFT calculation describes well the crossover behavior of Ce $4f$ electrons with one $T_0^F(\sim130$ K) for the participation of $4f$ electrons in the conduction and another $T_m^0(\sim50$ K) for the formation of coherent heavy electron $4f$ bands. The increase of the scattering rate approaching $T_m^0$ from high $T$ can be comprehended by the local Kondo effect scaled by $T_0^F$. The decrease from the maximum scattering rate with lowering $T$ is understood by the lattice coherence, which is consistent with the meaning of $T_m^0$.

We have examined the evolution of the heavy fermion state using electronic structure methods. As shown in the two-fluid phenomenology [14], the experimental studies of other heavy fermion systems [31] as well as the slave boson studies [32,33], the crossover from the high $T$ regime, where moments and quasiparticles coexist, to the low $T$ Fermi liquid heavy fermion state, has a rich structure characterized by multiple energy scales. We have found that it is characterized by multiple scales which have a clear correspondence with physical observables. $T_{m}^0$ is the onset of the sharp crossover where the small FS begins distorting towards the low $T$ FS. At a lower $T_{m}^0$, composite quasiparticles formed from $f$ moments and conduction electrons emerge, and this is signaled by a maximum of the resistivity. By that point, the FS has reached a shape which is closer to its zero temperature final value, but the material is not yet a Fermi liquid, which is only reached at a much lower temperature $T_{FL}$. We can only put bounds for this quantity as being lower than 10 K for the 115 material.

The theory can be tested using several techniques such as ARPES, Compton scattering, and scanning tunnelling microscopy, which have been developed as powerful tools for exploring the evolution of the electronic structure and are currently under way [34]. Our theory predicts that both $T_{m}^0$ and $T_0^F$ increase as a function of pressure in the CeIrIn₅ material. More generally, it would be interesting to follow these scales as a function of control parameters such as pressure and composition, to investigate the behavior of $T_{m}^0$ and $T_0^F$ in related materials which can be driven to a QCP.

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[34] J.D. Delinger et al. (private communication).