Electronic structure and the Fermi surface of UTGa$_5$
($T=$Fe, Co, Rh)

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Abstract

The relativistic energy-band calculations have been carried out for UFeGa$_5$, UCoGa$_5$ and URhGa$_5$ under the assumption that $5f$-electrons are itinerant. A hybridization between the U $5f$ state and Ga $4p$ state occurs in the vicinity of the Fermi level. The Fermi surface of UCoGa$_5$ is quite similar to that of URhGa$_5$, which are all small in size and closed in topology. UFeGa$_5$ has the quasi-two-dimensional Fermi surface which looks like a lattice structure.

Key words: Relativistic APW method; Fermi surface; actinide compound

Uranium compounds exhibit a great variety of phenomena such as Pauli paramagnetism, quadrupolar and magnetic ordering, heavy fermion, and unconventional superconductivity. It is considered that these behaviours are closely related to the crystal structure, correlation among $f$-electrons, and the $f$-electron hybridization with conduction and/or other valence electrons. To understand the electronic properties of these materials, the features of the ground state should be first clarified. For this purpose, it is quite important to investigate the Fermi surface both experimentally and theoretically.

UTGa$_5$($T$:transition atom) belongs to the heavy fermion compounds with the HoCoGa$_5$-type tetragonal crystal structure. In order to calculate the energy band structure of uranium compounds, relativity should be taken into account because of large atomic number. In this paper, the energy band structures for UFeGa$_5$, UCoGa$_5$ and URhGa$_5$ are calculated by the relativistic linear augmented-plane-wave(RLAPW) method[1]. In the RLAPW method, the four-component relativistic plane wave is augmented by a linear combination of the solutions for the spherically symmetrical potential. The exchange and correlation potential is treated in the local density approximation(LDA). The spatial shape of the one-electron potential is determined in the muffin-tin approximation. The self-consistent calculation is carried out for the experimental lattice constant.

In Fig. 1, the energy band structure calculated for UCoGa$_5$ is shown. The Fermi energy $E_F$ is located at 0.460 Ryd. Narrow bands, which lie just above $E_F$ and are split into two subbands by the spin-orbit interaction, are identified as the U $5f$ bands. A hybridization between the U $5f$ state and Ga $4p$ state occurs in the vicinity of $E_F$. The 15th and 16th bands form the Fermi surface, which is shown in Fig. 2. These sheets of the Fermi surface are all small in size and closed in topology. The hole sheets of the Fermi surface, as shown in Fig. 2(a), consists of the sheet centered at the $\Gamma$ point, two equivalent sheets centered at the $X$ points and the sheet which lies across the $\Sigma$ axis. Fig. 2(b) shows a set of the sixteen electron sheets of the Fermi surface in the 16th band. Each electron sheet lies across the T axis and looks like a cushions. The total number of holes is equal to that of electrons, which means that UCoGa$_5$ is a compensated metal.

The theoretical electronic specific-heat coefficient

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γ_{band} is 4.2 mJ/K^2·mol. It is about the half of the experimental value of 7 mJ/K^2·mol [2]. This disagreement between γ_{band} and γ_{exp} should be ascribed to the effects of the electron correlation which the LDA fails to take into account.

Similarly, we calculated the energy band structure of URhGa₅ with the RLAPW method in the LDA. The 15th and 16th bands construct the Fermi surface, which are very similar to those of UCoGa₅. Fig. 3 shows the total density of states as a function of energy in the vicinity of E_F.

Finally, the energy band structure of UFeGa₅ is discussed. The 14th and 15th bands construct the Fermi surface sheets. UFeGa₅ has the odd number of electrons per primitive cell, and therefore becomes the uncompensated metal where the numbers of electrons and holes on the fermi surface are not equal to each other. The electron sheets of the Fermi surface in the 15th band are cylindrical, which look like the lattice structure. The origins of the de Haas-van Alphen frequency branches can be clarified satisfactorily well by our theoretical fermi surface model[3]. Note that the electron sheets are quite analogous to those of CeIrIn₅[4]. Due to the simple two-dimensionality of the Fermi surface, CeIrIn₅ is an ideal example for fundamental study of superconductivity in rare earth compounds. At present, we construct a microscopic model by combining the band calculation and the tight-binding method[5,6].

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