Non-fermi liquid behaviour in uranium-based heavy-fermion compounds

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8. Concluding remarks

8.1. $U_2Pt_2In$

In this thesis, it has been shown that $U_2Pt_2In$ is the first stoichiometric U-based compound that exhibits non-Fermi liquid (NFL) behaviour at ambient pressure. Research on stoichiometric compounds prepared in a single-crystalline form is highly desirable in the study of NFL behaviour because the effects of disorder may not dominate the physics. This provided a strong motivation to carry out an extensive study on $U_2Pt_2In$ in order to determine its NFL properties. Nevertheless, the rather difficult metallurgy of $U_2Pt_2In$ imposed its restrictions on the research carried out.

The main results obtained on $U_2Pt_2In$ are summarized below.

- Polymorphism was found in $U_2Pt_2In$ [1]: while polycrystalline samples form in the $U_3Si_2$-type of structure, single-crystals form in the $Zr_3Al_2$-type of structure (superstructure of the $U_3Si_2$ type with a doubling along the c-axis).
- When tracing the magnetic ordering temperatures of the $U_2T_2X$ family of compounds versus the square of the conduction-electron - $f$-electron hybridization matrix element in a Doniach-like diagram, one finds that $U_2Pt_2In$ is located at the border line between magnetic and non-magnetic compounds, which suggests that $U_2Pt_2In$ is near a magnetic instability.
The magnetic susceptibility of U$_2$Pt$_2$In is weakly anisotropic and follows a (modified) Curie-Weiss behaviour at high temperatures. At low temperatures, however, an anomalous behaviour is observed: $\chi$ goes through a maximum at 7.9 K, which is attributed to the stabilization of short-range antiferromagnetic (AF) correlations, while $\chi_\parallel$ increases as $1-bT^{-0.7}$ for $2 \text{ K} \leq T < 10 \text{ K}$. This is at variance with the standard Fermi liquid (FL) behaviour, which predicts $\chi$ to attain a constant value at low temperatures.

The resistivity of U$_2$Pt$_2$In is highly anisotropic below about 80 K with $\rho_\perp > \rho_\parallel$. At the lowest temperatures, the resistivity does not follow the FL quadratic temperature dependence, instead $\rho_\parallel \sim T^{1.25}$ and $\rho_\perp \sim T^{0.9}$. The residual resistivities are unusually high: $\rho_{0,\parallel} = 115 \mu\Omega\text{cm}$ and $\rho_{0,\perp} = 210 \mu\Omega\text{cm}$. However, structure refinements from X-ray [1] and neutron-diffraction [2] experiments indicate a high sample quality. Also, the large difference between $\rho_{0,\parallel}$ and $\rho_{0,\perp}$ shows that $\rho_0$ is largely determined by other scattering mechanisms than impurity or defect scattering. It should also be noted that the higher resistivity values are found for a current along the e-axis, which is the axis along which the AF fluctuations stabilize.

Magnetoresistance (MR) experiments show a gradual increase of the resistivity exponent $\alpha$ (defined as $\rho \sim T^\alpha$ as $T \to 0$) with increasing magnetic field strength. At 8 T, $\alpha = 2$ as expected for a FL. There are two different contributions to the MR: a negative one (associated with spin effects) and a positive one (associated with orbital effects). At low fields, the negative contribution is dominant, except in the case $B \parallel I \parallel a$, whereas at high fields the positive contribution becomes more and more important. The negative contribution to the MR is more important for $I \parallel e$ than for $I \parallel a$. However, no satisfactory explanation for this anisotropy can be offered at the moment. Additional evidence that the high residual resistivity in U$_2$Pt$_2$In is not determined by crystallographic disorder is provided by the strong field dependence of $\rho_0$.

The specific heat of U$_2$Pt$_2$In provides solid evidence for the classification of this heavy-fermion compound as a NFL. The specific heat shows a pronounced diverging behaviour of the type $c/T \sim -\ln(T/T_0)$ over almost two decades of temperature. The low-temperature specific heat is dominated by a strong contribution of the In nuclear moments. The same logarithmic divergency is observed in the specific heat measured on polycrystalline samples. As these crystallize in the simpler U$_2$Si$_2$-type of structure, the NFL behaviour is not directly related to the presence of two inequivalent U positions, as present in the single crystals with the Zr$_3$Al$_2$-type of structure.
• The thermal-expansion coefficient of U$_2$Pt$_2$In becomes anisotropic below about 12 K, in a way that the c-axis shrinks more rapidly than the a-axis upon formation of the heavy-fermion bands.

• Absence of weak magnetic order, at least down to 0.05 K, is confirmed by means of muon spin relaxation and rotation (μSR) experiments. Besides a static magnetic component originating from the In nuclear moments, the μSR spectra below 10 K reveal the presence of magnetic fluctuations. No evidence was found for Kondo disorder.

• Resistivity experiments under hydrostatic pressure indicate a recovery of the FL $\rho \sim T^2$ behaviour at low temperatures for $I \parallel a$. This is consistent with predictions from a transport theory for heavy-fermion compounds near an AF quantum critical point (QCP) [3]. The anisotropy in the resistivity is strongly enhanced under pressure, as follows from the increase of the ratio $\rho_c/\rho_a$ measured for 0.3 K $\leq T < 300$ K. Due to the enhancement of the anisotropy, the resistivity curves for $I \parallel c$ do not show a $\rho \sim T^2$ behaviour but a low-temperature minimum.

One of the main issues that arise when discussing the properties of NFL compounds is the responsible mechanism for this behaviour. Although in the current state-of-the-art NFL physics no definite answers can be provided, one most important distinction can often be made: whether the NFL behaviour is due to a single-ion or a cooperative mechanism.

The location of U$_2$Pt$_2$In at the border line between magnetic and non-magnetic compounds in the Doniach diagram for the U$_2$T$_2$In family of compounds, suggests a proximity to an AF QCP. Resistivity measurements under pressure carried out on U$_2$Pt$_2$In and U$_2$Pd$_2$In yield strong support for this hypothesis. Considering the absence of magnetic order down to 0.05 K (as evidenced from the μSR experiments) and the important finding that pressure leads to the recovery of the FL behaviour, one cannot exclude that U$_2$Pt$_2$In is even located at the AF QCP. In order to investigate this further, specific-heat experiments under pressure would be most welcome. As the specific-heat coefficient $\gamma$ is related to the coherence temperature $T_{coh}$ [4], the observation and evolution of the FL $\gamma$ coefficient with pressure would provide valuable information on the recovery of the FL state near the QCP.

It is important to notice that the observed divergency of the specific heat, $c/T \sim -\ln(T/T_0)$ is not consistent with an AF QCP, but rather indicates a ferromagnetic (FM) QCP [5]. However, a logarithmic divergency of $c/T$ is allowed for an AF QCP in a two-dimensional (2D) system. This appears to apply to compounds like CeCu$_{50}$Au$_{50}$ [6] and CeNi$_2$Ge$_2$ [7], where inelastic neutron
scattering (INS) experiments have provided evidence for a spectrum of strongly anisotropic magnetic fluctuations, with a quasi-2D nature. INS experiments on U\textsubscript{2}Pt\textsubscript{2}In could possibly elucidate the nature of the magnetic fluctuations further.

In order to investigate the evolution of magnetic order near U\textsubscript{2}Pt\textsubscript{2}In in the Doniach phase diagram, the study of single-phase samples of e.g. (U\textsubscript{1-x}Th\textsubscript{x})\textsubscript{2}Pt\textsubscript{2}In and U\textsubscript{2}(Pt\textsubscript{1-x}Pd\textsubscript{x})\textsubscript{2}In is helpful. If Pd or Th doping results in the emergence of magnetism, strong evidence for U\textsubscript{2}Pt\textsubscript{2}In being at a QCP is obtained. One could then also study such doped samples under pressure, which should lead to the suppression of magnetic order and to the appearance of the FL state.

The origin of the strongly anisotropic character of the resistivity of U\textsubscript{2}Pt\textsubscript{2}In remains puzzling. Its enhancement under pressure indicates that the anisotropy is not inherent to the crystal structure since the compressibility is almost isotropic. Resistivity measurements on single crystals of the non-magnetic compound Th\textsubscript{2}Pt\textsubscript{2}In would be useful to address this issue. These could also serve to obtain an estimate for the phonon contribution to the resistivity of U\textsubscript{2}Pt\textsubscript{2}In. In addition, specific-heat experiments on Th\textsubscript{2}Pt\textsubscript{2}In should be carried out in order to estimate the phonon contribution to \(\alpha(T)\). This would enable a more precise determination of the temperature up to which a logarithmic divergency is present in the electronic specific heat of U\textsubscript{2}Pt\textsubscript{2}In.

The high residual resistivity of U\textsubscript{2}Pt\textsubscript{2}In raises the question of the role of disorder in the NFL properties. The dependence of \(\rho_0\) on the current direction, as well as its field and pressure variations, indicate that impurity and defect scattering are not the dominant mechanisms leading to the high values of \(\rho_0\). Even though X-ray and neutron-diffraction experiments indicate a good single-crystalline quality, the use of a local probe to measure the near-neighbour bond-length distributions, like (synchrotron radiation) X-ray absorption fine-structure (XAFS) experiments, could be helpful to determine the exact amount of disorder. For instance, neutron-diffraction experiments carried out on UCu\textsubscript{4}Pd were inconclusive regarding the presence of disorder [8]. However, XAFS experiments on the same sample revealed the presence of Pd/Cu site interchange [9]. It was concluded that the amount of disorder observed was sufficient for the Kondo disorder model to apply.

Although there is no evidence for Kondo disorder in U\textsubscript{2}Pt\textsubscript{2}In, a good test for the Kondo disorder model is obtained by a comparison of the \(\mu\)SR and NMR line widths [10]. As shown in this work, the small frequency shifts observed in the transverse-field (TF) \(\mu\)SR spectra of U\textsubscript{2}Pt\textsubscript{2}In, require that a full analysis of the TF line widths can only be accomplished with samples with a well-defined geometry in order to account for demagnetizing effects.
Concluding remarks

A further study of the NFL behaviour of U3PtIn may hopefully be carried out on a "second generation" of single crystals, with a minimum amount of disorder. Single crystals with a lower residual resistivity should be prepared. Therefore, one should look into single-crystal growth methods other than the mineralization technique, by which possibly crystals of higher quality can be produced.

8.2. Related compounds

The presence of a QCP has also been investigated for the compounds U3Ni3Sn4 and U(Pt1-xPd,x)3.

Specific-heat measurements carried out on the stoichiometric compound U3Ni3Sn4 show the presence of a NFL regime in the temperature range 0.5-5 K and a crossover to a FL ground state below 0.4 K. The divergency of the specific heat in the NFL regime is of the type $c/T \sim \gamma_0 - \alpha T^{1/2}$, which is consistent with the proximity to an AF QCP in a 3D system [5]. Both the NFL $c/T$ divergence and the spin-fluctuation term in the FL specific heat, have the same characteristic temperature $T_0$ of 10 K. The pressure dependence of the temperature below which the FL regime in the resistivity is attained, is well described in terms of a transport theory for nearly AF metals [3]. These results indicate that by a small lattice expansion, equivalent to a negative pressure of about -0.04 GPa, U3Ni3Sn4 may be tuned to the QCP. The isostructural compound U3Cu3Sn4 is an antiferromagnet with $T_N = 13$ K and its unit-cell volume is about 5% larger than that of U3Ni3Sn4. Therefore, it would be highly interesting to check the existence of an AF QCP by studying samples in which small amounts of Ni are replaced by Cu.

μSR experiments on the system U3(Pt1-xPd,x)3 indicate a new type of QCP in the phase diagram: at the critical concentration $x_c = 0.006$, unconventional superconductivity is suppressed and a large-moment antiferromagnetic (LMAF) phase emerges. The fact that the superconducting wave-function has odd parity suggests that doping with Pd leads to a shift of the spectral weight from FM to AF fluctuations. The observed competition between superconductivity mediated by FM fluctuations and static AF order, is in contrast with superconductivity mediated by AF interactions in materials like CePd2Si2 close to the QCP [11]. The phase diagram of U(Pt1-xPd,x)3
is similar to the phase diagrams observed for high-temperature superconductors with the peculiarity that the critical points $T_N \to 0$ and $T_c \to 0$ coincide.

Resistivity measurements carried out on $U(Pt_{1-x}Pd_x)$ show values of the exponent $\alpha$ equal to 1.8 and 1.6 for samples with $x = 0.004$ and $x = 0.007$, respectively [12]. These values are inconclusive with respect to the type of QCP in the system ($\alpha = 3/2$ and $\alpha = 5/3$ are predicted for the AF and FM QCP, respectively). A systematic study of the resistivity exponents in samples with $x$ around 0.006 is required to clearly determine the type of magnetic fluctuations at the QCP.

The origin of the NFL behaviour observed in $URh_{1/3}Ni_{2/3}Al$ might be of the single-ion type. $URh_{1/3}Ni_{2/3}Al$ shows a diverging specific heat of the type $c/T \sim -\ln(T/T_0)$ below about 6 K. Below 10 K the resistivity increases as $\rho \sim 1-\alpha T^\alpha$ with $\alpha$ about 1. The low-temperature resistivity increase is an indication that the mechanism responsible for the NFL behaviour is of the single-ion type. In fact, a loss of coherence is predicted to occur at low temperatures within the Kondo disorder model. This model also predicts a logarithmic divergency of $c/T$ and the linear increase of the resistivity below $T_K$ as observed for $URh_{1/3}Ni_{2/3}Al$. A detailed structural analysis on single-phase samples is required to establish the amount of disorder in this compound.

References