Strong in-plane anisotropy in the electronic structure of fixed-valence $\beta$-LuAlB$_4$

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The origin of intrinsic quantum criticality in the heavy-fermion superconductor $\beta$-YbAlB$_4$ has been attributed to strong Yb valence fluctuations and its peculiar crystal structure. Here, we assess these contributions individually by studying the isostructural but fixed-valence compound $\beta$-LuAlB$_4$. Quantum oscillation measurements and DFT calculations reveal a Fermi surface in $\beta$-LuAlB$_4$ markedly different from that of $\beta$-YbAlB$_4$, consistent with a ‘large’ Fermi surface in the latter. We also find an unexpected in-plane anisotropy of the electronic structure, in contrast to the isotropic Kondo hybridization in $\beta$-YbAlB$_4$.

At a quantum critical point (QCP), a continuous, zero-temperature phase transition occurs when the ordered and disordered phases are energetically degenerate. Usually, this requires tuning materials to some material-dependent critical pressure, composition or magnetic field [1]. However, there are rare cases where quantum critical phenomena can be observed without tuning. Studying such ‘intrinsic’ quantum criticality aims at identifying robust mechanisms for unusual electronic phases, raising the prospect of unconventional superconductivity [2, 3].

The heavy-fermion superconductor $\beta$-YbAlB$_4$ is an example for such an intrinsically quantum critical system [4]. Initial evidence for its non-accidental nature was the observation of quantum critical behavior in this stoichiometric, clean system at zero magnetic field, which was later extended to finite pressures, providing an example for a quantum critical phase [5, 6]. Transport and thermodynamic measurements identified non-Fermi liquid behavior, yet the Wiedemann-Franz law is obeyed, demonstrating that quasiparticles remain intact [4, 7, 8].

It was realised that the peculiar crystal environment of the Yb atoms, as well as strong Yb valence fluctuations are central for understanding the physics of $\beta$-YbAlB$_4$ [9–11]. The material crystallizes into the orthorhombic ThMoB$_4$ structure where a distorted honeycomb lattice of Yb and Al atoms is sandwiched between layers of B atoms. As shown in Fig. 1, the B atoms occupy independent sites B1–B3, which give rise to a network of five- and sevenfold boron rings [12]. The Kondo hybridization induced by the orbital overlap within the Yb-B-ring structure was argued to be strongly anisotropic between the crystal $c$ axis and the $(ab)$ plane, but isotropic within the $(ab)$ plane [13]. This hybridization can induce the observed $T/B$ scaling of the magnetic susceptibility, and can also explain the intrinsic quantum criticality in $\beta$-YbAlB$_4$ [5, 13–18]. Recently, strong evidence supporting the in-plane isotropy was found in the linear dichroism of core-level x-ray photoemission and NMR studies [19, 20].

The isomorph $\alpha$-YbAlB$_4$ features the same local symmetry and similar valence fluctuations of the Yb atoms, as well as an essentially identical in-plane isotropy of the Kondo hybridization [9, 12, 19, 21]. However, it does not show superconductivity nor intrinsic quantum criticality [12]. This indicates that the effect of the lattice cannot be truncated to the local environment of the Yb atoms, and/or that subtle differences in the Yb-B structures between $\alpha$- and $\beta$-YbAlB$_4$ are pivotal [20, 21].

It is therefore paramount to assess the importance of the lattice structure in detail [22, 23]. For this purpose, $\beta$-LuAlB$_4$ is ideally suited, as it crystallizes into the same structure as $\beta$-YbAlB$_4$, with nearly identical lattice parameters [12, 24]. The Lu 4$f$ states are fully occupied and fall well below the Fermi level. Thus, the electronic structure of $\beta$-LuAlB$_4$ represents the ‘small’ Fermi surface limit of $\beta$-YbAlB$_4$, and the Kondo effect and valence fluctuations are suppressed.

In this Rapid Communication, we investigate the electronic structure of $\beta$-LuAlB$_4$ using Shubnikov-de Haas oscillation measurements. We resolve all Fermi surface sheets predicted by DFT calculations and we extract the charge carrier masses. Our findings are broadly consistent with DFT predictions, validating a closer look at the electronic structure in real space. This reveals that the Lu-B hybridization and the charge distribution in the unit cell are strongly anisotropic, in contrast to the isotropic Kondo hybridization in $\beta$-YbAlB$_4$.

Using the Al flux method [12], thin ($d \lesssim 10\mu$m), plate-like single crystals of $\beta$-LuAlB$_4$ were grown, with the crystal $c$ axis normal to the plates, confirmed by X-ray diffraction measurements. For the resistivity mea-
to (non-critical) strongly-correlated systems, where electron-electron scattering leads to a Fermi liquid $T^2$ behavior. The $T^5$ dependence is an indicator for dominant quasi-elastic electron-phonon scattering instead.

In order to determine the electronic structure in detail, we performed Shubnikov-de Haas measurements up to $\mu_0 H = 18$ T as a function of temperature and field orientation. A selection of magnetoresistivity curves is shown in Fig. 1(b) which demonstrate that the magnetoresistivity is essentially temperature-independent over more than two decades up to $T \approx 9$ K. This observation suggests that quasiparticle mobilities and densities remain constant in this temperature regime, which is consistent with the effectively constant zero-field resistivity below $T \approx 10$ K, Fig. 1(a). However, this observation is in marked contrast to $\beta$-YbAlB$_4$ where a nearly 80 % decrease in mobilities was found below 10 K upon approaching the quantum critical point [31].

From Fig. 1(b), and in particular for $H \parallel (100)$, a strong oscillatory component of the high-field resistivity can be seen [30]. On one hand, the large amplitudes of these quantum oscillations demonstrate the high quality of the sample studied. On the other hand, the oscillations remain visible even for $T = 9$ K, suggesting small quasiparticle masses $m$, equivalent to large mobilities $|\mu| \sim 1/m$.

We now turn to a quantitative analysis of these quantum oscillations. In Fig. 1(c), a smooth, low-order polynomial background has been removed. When plotted against $1/H$, multiple frequencies $f$ are apparent, which correspond to extremal cross-sectional areas $A$ of the Fermi surface perpendicular to the magnetic field [30] $f = Ah/(2\pi e)$. We employ an FFT, shown in Fig. 2(a), to extract the oscillation frequencies and amplitudes as a function of temperature and field orientation (see also the Supplemental Material (SM) [29]). All obtained frequencies are shown in Fig. 2(b) on top of the FFT spectra, represented as a background color scale. We identify a set of branches for fields close to the (010) direc-
tion, labelled $\beta_{3-6}$. These branches display an approximate $1/\sin(\theta)$ dependence, where $\theta$ measures the angle away from (001). Such a dependence is consistent with a weakly warped two-dimensional Fermi surface pocket along the reciprocal $b$ axis. Next, we find two weakly orientation-dependent lines for $3.4 \, \text{KT} \leq f \leq 4.5 \, \text{KT}$ and $f \geq 6.8 \, \text{KT}$ which we call $\delta$ and $2\delta$. Their flat dependencies are consistent with nearly spherical Fermi surface pockets, however the integer ratio between their frequencies suggests that $2\delta$ is the first harmonic of $\delta$. Finally, multiple frequencies below $0.9 \, \text{KT}$ are found with no obvious analytical dependence, labelled $\alpha$, $\gamma$, $\epsilon$ and $\beta_7$.

Next, cyclotron masses are extracted by following the oscillation amplitudes as a function of temperature, as shown in Fig. 2(c). Fitting the Lifshitz-Kosevich form [30], we find $m \approx 0.29m_e$ for $\alpha$, which is much smaller than any cyclotron mass reported on the more strongly correlated $\beta$-YbAlB$_4$ [32]. The integer ratios between the frequencies and masses of orbits $\alpha$, $2\alpha$ and $3\alpha$ allow us to identify $\alpha$ as a fundamental frequency, and $2\alpha$ and $3\alpha$ as harmonics. Further orbits are discussed in the SM [29]. For the following discussion, we will omit harmonics identified in this way, as summarised in Fig. 3(c).

To better understand the observed quantum oscillations, we now turn to our DFT calculations. In Fig. 3(a), we show the computed band structure and density of states (DOS) over a wide energy range. The localized Lu $4f$ states are easily identified about $4.1 \, \text{eV}$ and $5.6 \, \text{eV}$ below the Fermi level $E_F$. In the vicinity of $E_F$, the DOS is dominated by Lu and B states whereas the Al states play a minor role. Two bands cross the Fermi level, which give rise to two Fermi surface sheets with equal volume, one hole-like and one electron-like, as shown in Fig. 3(b). The hole-like sheet consists of two strongly warped cylinders along (010) which give rise to the extremal orbits $\beta_{1-8}$. They are joined by two three-dimensional pockets, producing orbits $\alpha_{1-3}$. The electron-like sheet consists of nearly spherical pockets in the corners of the first Brillouin zone which are the origin of the orbit $\delta$. Further pockets with varying sizes surround the $\Gamma$ point, with orbits labelled $\gamma_{1-3}$ and $\epsilon$.

In Fig. 3(c), we compare the computed extremal Fermi surface cross-sections translated into quantum oscillation frequencies to the experimentally determined fundamental frequencies. The agreement is convincing: the line $\delta$ and the branches $\beta_{3-6}$ are reproduced, and most frequencies below $0.9 \, \text{KT}$ can be mapped within experimental resolution ($\Delta f \approx 40 - 100 \, \text{KT}$ FWHM depending on field range). DFT slightly underestimates the size of the large Fermi surface pockets, i.e. predicted frequencies are slightly smaller than observed ones, as evident from Fig. 3(c). This applies to both electron-like and hole-like bands, consistent with an overall charge balance.

Having determined the electronic structure of $\beta$-LuAlB$_4$, the isostructural but fixed-valence reference compound to $\beta$-YbAlB$_4$, we now discuss the two main implications of the findings. Firstly, the frequencies of the quantum oscillations and their angular dependence, as well as the cyclotron masses reported here differ greatly from results on $\beta$-YbAlB$_4$ studied previously [32]. The most notable difference is the absence of the spherical pocket $\delta$ in $\beta$-YbAlB$_4$. Conversely, large cross-
sectional areas were observed in $\beta$-YbAlB$_4$ (dubbed $\beta$ in Ref. 32), which have no counterpart in $\beta$-LuAlB$_4$. Such substantial differences in the Fermi surface suggest that in the presence of large magnetic fields, the Fermi surface of $\beta$-YbAlB$_4$ is not ‘small’ but ‘large’ and the 4f hole is delocalized [32]. Moreover, the experimental cyclotron masses of $\beta$-LuAlB$_4$ are in good agreement with predicted values (Table SMI [29]), but they are on average a factor 10 smaller than in $\beta$-YbAlB$_4$ [32]. This reflects the significant mass enhancement due to the presence of correlated f-states near the Fermi energy.

The second implication is most evident from the hole-like Fermi surface sheet of $\beta$-LuAlB$_4$, Fig. 3(b): There is a strong anisotropy of the electronic structure within the (ab) plane. To trace its origin, we reiterate that the DOS at the Fermi level $E_F$ has predominant Lu and B character, as shown in Fig. 3(a). It is therefore suggestive to attribute the anisotropy to the Lu-B hybridization. This is confirmed when returning to real space: in Fig. 4(a) and (b), we show the computed charge distribution in the unit cell for states close to $E_F$. We find a much larger orbital overlap between the Lu $d_{z^2}$ and the B3 $p_z$ states, when compared to B1 and B2. A similar result is obtained when comparing the contributions of the B atoms to the DOS, Fig. 4(c). Over a wide energy range around $E_F$, the B3 states contribute roughly 30% more than B1 and B2, which originates fully from their $p_z$ orbitals.

This ineqivalence of the B atoms cannot be assigned to the form of the aforementioned atomic orbitals, since B $p_z$ and Lu $d_{z^2}$ orbitals are rotationally invariant along the crystal c axis. Hence, no anisotropy is expected for an isolated stack of Lu atoms and ideal B rings. Consequently, we must link it to the variation in Lu-B bond lengths (where the Lu-B3 bond is the shortest), and/or the global charge distribution within the orthorhombic unit cell. As shown in Fig. 4(b), the charge carrier density is indeed strongly anisotropic: it is the largest within the Lu-B3-B3-Lu structure along the crystal b axis, whereas the B1 and B2 atoms can be projected onto low-density stripes. The Al sites are effectively disconnected from charge carriers occupying states close to $E_F$ which marks them as ideally suited for chemical doping minimizing quasiparticle scattering.

We conclude by discussing the implications on $\beta$-YbAlB$_4$. As it is isostuctural to $\beta$-LuAlB$_4$, and since the Lu/Yb-B bond length variations are nearly identical (4.0% in critical $\beta$-YbAlB$_4$ and 3.8% in non-critical $\beta$-LuAlB$_4$, in contrast to 5.1% in non-critical $\alpha$-YbAlB$_4$ [12]), we would expect that a similarly anisotropic orbital overlap should prevail in $\beta$-YbAlB$_4$ (see also the SM for a detailed comparison [29]). However, previous studies in quantum critical $\alpha$- and $\beta$-YbAlB$_4$ found a Kondo hybridization which is isotropic in the (ab) plane [19, 20]. Our finding therefore provides evidence that the isotropic Kondo hybridization and intrinsic quantum criticality in $\beta$-YbAlB$_4$ emerge from an anisotropic electronic structure [9, 10, 16, 21]. Identifying a mechanism which counteracts this electronic anisotropy and/or (self-)stabilizes the quantum critical phase remains an open problem.

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FIG. 4. Inequivalence of the boron atoms. (a) Charge carrier distribution $\rho_{el}$ in the proximity of the Lu atom and the sevenfold B ring. Electronic states within $\Delta E = k_B \cdot 300$ K of the Fermi level were included. The closed surfaces encapsulate regions with $\rho_{el} > 0.55 \cdot 10^{21}/\text{cm}^3$. (b) The charge carrier distribution in the Lu/Al layer of the unit cell, with the B atoms projected. (c) Comparison of the density of states contributions of the B atoms. Upper panel: total contribution, lower panel: $p_z$ states only.

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