Fermi Surface Modeling of Light-Rare-Earth Hexaborides using Positron Annihilation Spectroscopy

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2D angular correlation of the positron annihilation radiation (2D-ACAR) spectra are measured for LaB$_6$ along high-symmetry directions and compared with first-principles calculations based on density functional theory (DFT). This allows the modeling of the Fermi surface in terms of ellipsoid electron pockets centered at X-points elongated along the \( \Sigma \) axis (\( \Gamma \rightarrow M \) direction). The obtained structure is in agreement with quantum oscillation measurements and previous band structure calculations. For the isostructural topologically nontrivial SmB$_6$, the similar ellipsoids are connected through necks that have significantly smaller radii in the case of LaB$_6$. A theoretical analysis of the 2D-ACAR spectra is also conducted for CeB$_6$ including the on-site repulsion \( U \)-correction to the local density approximation (LDA + \( U \)) of the DFT. The similarities of the 2D-ACAR spectra and the Fermi surface projections of these two compounds allow to infer that both LaB$_6$ and CeB$_6$ are topologically trivial correlated metals.

1. Introduction

The rare-earth hexaborides (RB$_6$, \( R = \text{La, Ce, Pr, Nd, and Sm} \)) provide an interesting subject for experimental and theoretical studies because of a variety of features that can be ascribed to the particular role played by their 4f-electrons.[2,3] The unfilled 4f-states are a challenging problem for an accurate theoretical description of the electronic structure of rare-earth compounds. The 4f energy levels often overlap with the non-4f-wide bands, form narrow resonances, but nevertheless are frequently treated as core states. A proper description, however, requires inclusion of the 4f-states in valence bands where these states are also subjected to a strong on-site Coulomb repulsion. A simple theory that can capture this strong on-site Coulomb repulsion for 4f-orbitals at the mean-field level is the local density approximation (LDA + \( U \)) approach.[3] Another important ingredient in the physics of the rare-earth compounds is the presence of significant spin-orbit coupling, which fortunately can be included on an equal footing with strong on-site Coulomb interactions (the Hubbard \( U \)). The recent theoretical models beyond the local approximations, such as GW,[4] or…
dynamic electronic correlations such as dynamical mean field theory (DMFT),\textsuperscript{5,6} however, are difficult to use in case of multiorbitals (like 4f) and the strong spin–orbit coupling.

In the R\textsubscript{6} series of compounds, LaB\textsubscript{6} is considered as the reference non-f-electron system, CeB\textsubscript{6} is a Kondo lattice-heavy fermion system, whereas PrB\textsubscript{6} and NdB\textsubscript{6} are local-moment magnetically ordered metals. Among these compounds, SmB\textsubscript{6} is considered to be a “topological Kondo insulator”\textsuperscript{7} due to the narrow hybridization bandgap that opens at the Fermi level. In general, some of the 4f-bands of the rare-earth elements are located around the Fermi level (E\textsubscript{F}) and are crossed by the more dispersive 5d band. The finite spin–orbit coupling splits these crossings and opens a gap. The band-character inversion due to the spin–orbit coupling between the rare-earth 4f and opens a gap. The band-character inversion due to the spin–orbit coupling between the rare-earth 4f and opens a gap. The band-character inversion due to the spin–orbit coupling between the rare-earth 4f and opens a gap. Therefore, we believe that future studies with DMFT, which takes into account the full local correlations starting from itinerant (valence) 4f-states, will provide a suitable description of the electronic structure of the rare-earth hexaborides.

In the present study, we investigate the electronic properties of two members of the RB\textsubscript{6} family, namely LaB\textsubscript{6} and CeB\textsubscript{6}, with 2D angular correlation of annihilation radiation (2D-ACAR) spectroscopy and first-principles density functional theory (DFT) calculations.\textsuperscript{10–13} We analyze the radial anisotropies of the obtained data in the extended momentum space and the corresponding back-folded spectra in the crystal-momentum space to identify the different Fermi surface (FS) features. We also compare the experimentally measured and computed 2D-ACAR spectra for LaB\textsubscript{6}. The crystal structures of RB\textsubscript{6} belong to the simple cubic CsCl structure type (Pm\textsubscript{3}m), with lattice constant a = 4.1569Å and a = 4.1391Å, for LaB\textsubscript{6} and CeB\textsubscript{6}, respectively. The rare-earth atoms occupy the corner of the unit cell, corresponding to the 1a(0,0,0) Wyckoff site, whereas the B atoms are located at the octahedral sites in the body-centered position at the 6f (0.5,0,0,0,0,0) Wyckoff sites, where z is 0.2011 for CeB\textsubscript{6}\textsuperscript{14} and 0.1996 for LaB\textsubscript{6}\textsuperscript{15}.

Although the f-shell of La is empty while Ce has a single f-electron and CeB\textsubscript{6} exhibits a far richer phase diagram than LaB\textsubscript{6}, undergoing several different magnetic phase transitions, de Haas–Van Alphen (dHvA) measurements\textsuperscript{16} yielded similar FS features for both CeB\textsubscript{6} and LaB\textsubscript{6}. The FS of CeB\textsubscript{6} consists of large “ellipsoid” pockets around the X points of the Brillouin zone (BZ), whereas the states around the zone center (G-point) are shifted away by band-renormalization effects leading to a hole pocket.\textsuperscript{17} The first-principles calculations agree with the experimental results around the X-points but fail to capture the strongly renormalized electronic states around the Γ-point.\textsuperscript{17} The dHvA measurements\textsuperscript{16} at temperatures T < T\textsubscript{K} can be brought into accordance with a model in which f-electrons do not contribute to the Fermi volume. Above the Kondo temperature T > T\textsubscript{K}, in the paramagnetic phase, dHvA measurements and 2D-ACAR results were also in agreement in the case of CeB\textsubscript{6}.\textsuperscript{16} Results of the 2D-ACAR experiments on LaB\textsubscript{6} were reported earlier in the literature,\textsuperscript{18,19} where the analysis in terms of the back-folded momentum densities were conducted. 3D reconstruction of the FSs using 2D-ACAR projections was also reported\textsuperscript{20} and contrary to the electronic structure calculations, the strong hybridization between B-p and La-d states (f-states are unoccupied) that produced additional FS sheets was not detected in the experiment.

In our work, we analyze the momentum densities of LaB\textsubscript{6} (experiment/theory) both in the extended p and in the reduced k spaces. Possible similarities between the theoretical spectra with the isostructural CeB\textsubscript{6} are also discussed. For the CeB\textsubscript{6} compound, in the absence of recent experimental measurements, we theoretically study the effects of the different U values of the Hubbard interaction and diverse double-counting schemes of the LDA+U method. Based on our 2D-ACAR results, we conclude that both LaB\textsubscript{6} and CeB\textsubscript{6} are topologically trivial but correlated metals. Therefore, we believe that future studies with DMFT, which takes into account the full local correlations starting from itinerant (valence) 4f-states, will provide a suitable description of the electronic structure of the rare-earth hexaborides.

2. Methods

2.1. 2D-ACAR

2D-ACAR measurement is a powerful tool to investigate the bulk electronic structure.\textsuperscript{21,22} It is based on the annihilation of positrons with electrons of a sample leading to the emission of two γ-quanta in nearly antiparallel directions. The small angular deviation from collinearity is caused by the transverse component of the electron’s momentum. Hence, the coincident measurement of the annihilation quanta for many annihilation events yields a projection of the so-called two-photon momentum density (TPMD) ρ\textsuperscript{2γ} (p). This is usually computed as the Fourier transform of the product of positron wave function Ψ\textsuperscript{0} (x) (thermalized positron in its ground state) and electron wave function Ψ\textsuperscript{−} (x):

\[
ρ\textsuperscript{2γ} (p) ∝ \sum_{\mathbf{k}} n_j (\mathbf{k}) \left| \int dx e^{-ip\cdot x} \Psi\textsuperscript{0} (x) \Psi\textsuperscript{−} (x) \sqrt{\gamma (x)} \right|^2
\]  

(1)

The sum runs over all \( \mathbf{k} \) states in all bands, \( j \), with the occupation \( n_j (\mathbf{k}) \). The so-called “enhancement factor” \( \gamma (x)\textsuperscript{23} \) takes into account the electron–positron correlation. The 2D-ACAR spectrum \( ρ\textsubscript{ACAR} (p_x, p_y) \), the quantity which is actually accessible by an experiment, is a 2D projection of the 3D momentum–density distribution \( ρ\textsuperscript{2γ} (p) \) along a chosen (p\textsubscript{z}) axis:

\[
ρ\textsubscript{ACAR} (p_x, p_y) = \int \rho\textsuperscript{2γ} (p) dp_z
\]

(2)

The positron annihilation probes all electrons in the system. Filled bands, especially bands of core electrons, give a nearly isotropic distribution, which is superimposed by an anisotropic contribution, mainly produced by the electrons near the FS. This anisotropic \( ρ\textsubscript{aniso} (p_x, p_y) \) contribution is therefore one of the most interesting features of an ACAR spectrum \( ρ\textsubscript{ACAR} (p_x, p_y) \). It can be calculated by subtracting the isotropic contribution \( ρ\textsubscript{ACAR} (p_x, p_y) \):
\[ \rho_{\text{ano}}(p_x, p_y) = \rho_{\text{ACAR}}(p_x, p_y) - \rho_{\text{ACAR}}(0, 0) \]

where the isotropic contribution is the radial average
\[ \rho_{\text{ACAR}}(p_x, p_y) \equiv \rho_{\text{ACAR}}(p_x^2 + p_y^2)^{1/2} \] constructed from the original spectrum \( \rho_{\text{ACAR}}(p_x, p_y) \) by averaging over all data points in equidistant intervals \( [p_x, p_y + \Delta p_x] \) from the center.

The discontinuities in \( \rho^{\text{fi}}(p) \) correspond to the density jumps in the Fermi distribution and are used to identify the FS sheets. These discontinuities occur at \( p = k + G \), where \( k \) is the wave-vector in the first BZ and \( G \) is a reciprocal lattice vector, which can be folded back from the extended zone scheme \( p \)-space to the reduced zone scheme \( k \)-space by applying the so-called lock-crisp-west (LCW) theorem.[24]

### 2.2. Experiment Details

The ACAR experiments were conducted at the 2D-ACAR spectrometer at the Technische Universität München. The detector–detector distance was 17.5 m with the sample positioned exactly in the middle. This led to an angular resolution of \( \sigma_n = 0.538(1) \text{ mrad} \) and \( \sigma_v = 0.655(1) \text{ mrad} \) in the horizontal and vertical direction, respectively.[25] All measurements were carried out at 15 K to reduce the contribution of the thermal motion of the positron. The positrons were guided onto the sample by a magnetic field of 1.2 T at the sample position. For full details on the experimental setup, see the study by Ceeh et al.[26]

The LaB\(_6\) sample used in the 2D-ACAR experiment was a cuboid-shaped high-quality single crystal of \( 10 \times 5 \times 2 \text{ mm}^3 \) size. It was grown by the vertical crucible-free inductive melting method. The grown crystal was characterized by Laue backscattering, optical spectral analysis, X-ray diffraction, and density measurements. The surfaces of the plate were polished with diamond powder.

We recorded 2D-ACAR projections along the high-symmetry [110], [001], and [111] directions within the (110) plane. For each spectrum, more than 9 × 10\(^7\) events were collected. The raw data were divided by the so-called momentum sampling function (MSF), to account for the sensitivity of the instrument to detect photon pairs of the given projected momentum (resulting from the finite field of view of the Anger camera detectors and spatial inhomogeneities in their detection efficiency).[27] To calculate the MSF, the single events on both detectors were collected in parallel to the coincident events throughout the experiment. The convolution of those single spectra gave the MSF. Due to the crystal symmetry, certain projections had to show defined symmetries. In the measured data, certain pixels are equivalent (e.g., in case of a (001) high-symmetry projection, the fourfold rotation symmetry and the mirror symmetry lead to eight equivalent pixels). Due to experimental noise, however, they usually do not have equal intensity and therefore, the data is symmetrized, that is, averaged over the equivalent pixels, which hence increases the statistics per pixel.[28]

### 2.3. Electronic Structure Calculation and Enhancement Models

Electronic structure calculations were carried out using the ELK code,[29] which is an all-electron full-potential linearized augmented plane-wave (LAPW) code for determining the properties of crystalline solids. The valence electron configuration for the rare-earth atoms (La/Ce) was 6s6p5d4f, whereas the valence electrons of B were located in the 2s and 2p orbitals. The self-consistent calculations were carried out using the local density approximation (LDA) exchange-correlation functional parameterized by Perdew and Wang.[30] For the LDA+U, the rationally invariant formulation of the Coulomb interaction between the valence electrons was used[31] with different double-counting schemes.[31] Further details of the LDA+U calculations can be found in Section 3.2. In the self-consistent calculations, the summations were conducted using the tetrahedron method, with a \( 20 \times 20 \times 20 \text{ } k \)-mesh in the irreducible part of the BZ, and convergence was achieved in total energy with an accuracy of better than 10\(^{-5}\) Ry. For the detailed FS calculations, a significantly larger, \( 80 \times 80 \times 80 \text{ } k \)-mesh was used.

The DFT can be generalized to electron–positron systems by including the positron density, in the form of the two-component DFT.[32,33] In this formalism, the positron was considered to be thermalized and described by a state with \( p_+ = 0 \) with \( s \)-type symmetry at the bottom of the positronic band. The photons resulting from the electron–positron annihilation carried the momentum of the annihilated pair, up to a reciprocal lattice vector, reflecting the fact that the annihilation took place in a crystal. Hence, an electron with a transverse wavevector \( k \) contributes to spin-resolved TPMD, \( \rho^{\text{fi}}(p) \), not only at \( p = k \) (normal process) but also at \( p = k + G \), with \( G \) being a vector of the reciprocal lattice (Umklapp process). In the LDA(+U) framework, the electron–positron momentum density \( \rho(p) \) was computed directly with the spin-resolved versions of (1), (2). We calculated the 2D-ACAR spectra according to the method described in the study by Ernsting et al.[34] These TPMDs were calculated to a maximum momentum of 6 a.u. and the electron–positron correlations were taken into account within the \( \gamma(x) \) term, the so-called enhancement factor. Within the independent particle model (IPM), \( \gamma(x) = 1 \), whereas models of enhancement beyond IPM consider momentum and energy dependencies and a separate treatment of \( f \) and \( d \) states in comparison with the \( s \) and \( p \) states. These have been quantified via many different approximations.[35] In the present studies, we used the so-called Drummond parametrization.[36,37]

### 3. Results and Discussion

#### 3.1. LaB\(_6\): Experiment and Theory

To compare experimental and theoretical data on LaB\(_6\) in the \( p \)-space, we look at the radial anisotropy \( \rho_{\text{ano}}(p_x, p_y) \), which emphasizes the spectral contribution from electrons near the Fermi level. Figure 1 shows the radial anisotropy for all three measured projections (from left to right (110), (001), and (111)) in the upper half and the corresponding theoretical calculations in the lower half of the plots. The calculated results are for the momentum distributions with IPM (left) and Drummond enhancement (right). As the experimental data are symmetrized according to the crystal symmetry, we convolved the theoretical data with an isotropic Gaussian with an effective resolution of...
σ_{exp} = 0.60 \text{ mrad} calculated as the quadratic mean of the experimental resolutions given in Section 2.2.

Overall, we have very good agreement between theory and experiment. Dominant features seen in the theoretical calculations are also visible in the experimental data. Especially the fourfold and sixfold symmetric projections are well reproduced. Some minor differences can be seen in the \((110)\) projection. Particularly, the weight distribution in the region between \(-5\) and \(5\) mrad on both axes shows a significant departure between the theory and experiment. However, the high-intensity positive signal along \([110]\) direction at about \(8\) mrad is well reproduced. We attribute the better agreement between the theory and the experiment for the \((001)\) and \((111)\) projections to the higher-point symmetry of these directions. In all experimental projections, the anisotropy profiles are slightly stretched out within the plane. The similar behaviour is also observed when the Drummond enhancement is included in the computation as compared with the IPM results. However, the radial expansion of the profile is not sufficient to match the observed “stretching” in the experimental measurement. The differences between the IPM and Drummond model are marginal, though.

Figure 2 shows the LCW back-folded data, experiments as well as theory, and is organized according to the structure of Figure 1. The top part of each plot shows the \(k\)-space experimental 2D-ACAR spectrum, whereas the bottom part is split between the calculated results using IPM (left) and Drummond enhancement (right), broadened with the experimental resolution.

The measured and calculated LCW back-folded spectra show an overall good agreement. The \((001)\) and \((111)\) projections show the expected features of the FS topology (see Figure 3). In the \((111)\) projection, the sixfold rose structure around the \(\Gamma\)-point becomes apparent. As expected from the calculated FS, there is no density at the \(\Gamma\) - and \(R\)-point in the BZ. Note that one can not see the hole space around the \(\Gamma\)-point in the \((001)\)
projection, as it is covered by the “ellipsoids” along the [001] direction. While the theoretical calculations explicitly reproduce the “ellipsoid” structure in (110) projection, the experimental (110) projection shows stripes along [1T0] direction. The latter can be also attributed to the observed substantial discrepancies between the theory and experiment in the radial anisotropy (see Figure 1).

The FS of LaB$_6$ is shown in Figure 3 in the simple cubic BZ where the high-symmetry points are $\Gamma \equiv [0, 0, 0]$, $X \equiv [1/2, 0, 0]$, $M \equiv [1/2, 1/2, 0]$, and $R \equiv [1/2, 1/2, 1/2]$. The FS of LaB$_6$ consists of a set of equivalent “ellipsoids” centered at the $X$-points and connected by necks which intersect along $\Gamma - M$ direction.

In the panels (b) and (c) in Figure 3, we show a cross section in the $\Gamma - X - M$ plane ($k_{[001]} = 0$ plane). The DFT (LDA) FS calculation for SmB$_6$, with a downward shift of the Fermi energy, is presented in the study by Tan et al.\cite{38} and has a FS similar to that of LaB$_6$, as shown in Figure 3. The angular variations of the dHvA frequencies and the disappearance of oscillations in some angular regions have been recently discussed for both SmB$_6$ and LaB$_6$.\cite{38} These quantum oscillation frequencies identify the so-called $\alpha, \gamma, \epsilon$ branches associated with FS regions visible in the $\Gamma - X - M$ plane, see Figure 3. The $\alpha$-branches were associated with the ellipsoids centered at the $X$-point. As one can see, the nearest-neighbor ellipsoid FS pieces touch along the $\Gamma-M$ line and connect through the small distorted circular shape neck, building up a multiple-connected FS. Both the $\gamma$ and $\epsilon$-orbits are hole-like orbits and are centered around $M$-point and $\Gamma$-point, respectively. According to the dHvA frequencies the angular region for the hole-like $\gamma$- and $\epsilon$-orbits is significantly larger in SmB$_6$ than in LaB$_6$, which may be the cause of different physical properties of these two compounds. From our calculations we can compare linear dimensions of the FS features to both previous computations and the present experimental data. The corresponding values are 0.848, 0.668, and 0.624 for the $X-\Gamma$, $X-M$, and $X-R$ directions, respectively, as a fraction of the BZ size. Our results agree with the previous results reported in the literature.\cite{20}

To determine the FS linear dimension from the experimental data,\cite{20,39} taking into account a finite experimental resolution, we model the electron–momentum density with prolate ellipsoids with equatorial radius $R_{eq}$ and polar radius $R_{pl}$, centered at the $X$ points in reciprocal space and pointing toward the neighbouring $\Gamma$ points.

$$F(k; R_{eq}, R_{pl}) = \sum_G \sum_{b_i} \Theta\left(1 - \frac{|(k - G) \times b_i|^2}{R_{eq}^2 |b_i|^2} - \frac{|(k - G - b_i/2) \cdot b_i|^2}{R_{pl}^2 |b_i|^2}\right)$$

Here, $\Theta(\cdot)$ is the Heaviside step function and summation is over reciprocal lattice vectors $G$ and reciprocal primitive vectors $b_i \in \{b_{[100]}, b_{[010]}, b_{[001]}\}$. Adding a flat background TPMD due to fully occupied bands described by a third parameter $F_{backg}$, projecting along the given direction and convolving with the anisotropic Gaussian kernel that models the experimental resolution, we fit the resulting simulated projections to the correspondingly normalized LCW back-folded data for all measured projection directions by varying the three free parameters. We obtain $2R_{eq} = 0.662(2)$ and $2R_{pl} = 0.952(2)$ in units of the reciprocal lattice constant, where the reported uncertainties are statistical precision due to counting noise. We see that the equatorial diameter $2R_{eq}$ agrees well with the calculated $X-M$ length, whereas the length along the $X-\Gamma$ direction is overestimated. We think that this is due to positron wave-function and/or electron–positron correlation effects, which are neglected in the LCW theorem.\cite{24} While the equatorial diameter is directly accessible in particular in the $\{001\}$ projection, where the contrast between the filled ellipsoids and the background density is large, there is no projection direction that would afford an unstructured view on the polar termination of the ellipsoids. Thus, a larger band weight of the conduction and/or core bands toward $\Gamma$ would necessarily result in an underestimated $R_{pl}$. Also, with the current experimental resolution, we cannot detect possible deviations of the FS from the shape of a rotational ellipsoid in the vicinity of the poles or whether the $X-M-R$ cross section of the FS shows fourfold deviations from circular symmetry, as shown by the calculations (see Figure 3).

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**Figure 3.** a,d) FS in the first BZ from first-principle LDA calculations for LaB$_6$ a) and d) LDA+U calculation with $U = 6$ eV for CeB$_6$. b,c) $\Gamma - X - M$ cross section ($\{010\} - \{001\}$ cut) of FS for LaB$_6$ and CeB$_6$, respectively. On plots (c) and (d), the FS sheets of CeB$_6$ for two different spin projections are depicted in the left and the right half of the plots, in blue/orange and deep sky blue/red colors, respectively. The borders of the first BZ (gray lines) and high-symmetry points ($\Gamma, X, M, R$) are shown on all plots as well.
Among the most frequently used techniques to include short-range Coulomb interactions between the electrons in the framework of DFT are the self-interaction-corrected local-spin-density approximation (SIC-LSDA)\cite{40} and LDA+U\cite{3}. The LDA+U method is particularly useful as it is computationally less demanding and hence can be easily used for systems with unit cells consisting of a considerable number of atoms. Furthermore, its flexibility allows a systematic study of effects of the on-site repulsion $U$, on the ground-state properties such as equilibrium lattice parameter, magnetic moment, and—in the current work—2D-ACAR spectra. The orbital-dependent LDA+U functional used in the present work is given by\cite{3}

$$E_{\text{LDA}+U} = E_{\text{LDA}} + \frac{U}{2} \sum_{\sigma} [\text{tr} \rho^\sigma - \text{tr} (\rho^\sigma \rho^\sigma)]$$ \hspace{0.5cm} (5)

Here, $\rho^\sigma$ (not to be confused with momentum densities) is the density matrix for the $f$-states and $U$ and $J$ are the local Coulomb and exchange Hund’s parameters. The lack of a diagrammatic expansion of the DFT total energy makes it difficult to model the effect of the local Coulomb interactions beyond the effects already captured by the exchange-correlation functionals. To avoid the double counting of such effects, several schemes have been proposed for different kinds of materials. One of them is the mean-field approximation to the Hubbard correction, the so-called “fully localized” limit (FLL). In this scheme, each localized (e.g., atomic) orbital is either full or completely empty. This formulation of the double-counting term mimics an expansion of the electronic energy around the strongly localized limit and thus tends to work quite well for strongly correlated materials with localized orbitals. For other systems, such as metals or “weakly correlated” materials, the excessive stabilization of occupied states due to the “$+U$” potential may lead to unphysical results, for example, the enhancement of the Stoner factor.\cite{31} A different double-counting scheme, the “around mean-field” (AMF), was introduced in the study by Czyzyk et al.\cite{41} and further developed in the study by Petukhov et al.\cite{31}. In the present study, the AMF double-counting scheme failed to produce the correct position of the $4f$ bands; therefore, FLL double counting was used in all presented results.

The DFT band structure of CeB$_6$ produces the manifold of Ce-4$f$ orbitals in the close vicinity of the Fermi level. According to the experimental study of Neupane et al.\cite{17} they have to be located at around 2 eV in the conduction band. Therefore, for the FLL limit, a reasonable parameter for the effective Coulomb interaction would be $U = 4$ eV. The band structure and the density of states (not shown) in the ferromagnetic states are characterized by dispersive 5$d$- and flat 4$f$-bands along the $M-X-M$ and $X-F-X$ directions. The flat bands are purely built from Ce-4$f$ states. The dispersive 5$d$ band around the $X$-point are situated at 2 eV below $E_F$ and touches the $B-2p$ states at the $\Gamma$ point. The position of these bands agrees fairly well with experiments.\cite{17,42}

In Figure 4, we present results for the calculated radial anisotropy of the 2D-ACAR spectra compared with CeB$_6$. The electronic calculations for LaB$_6$ were done using DFT only. For CeB$_6$, we used LDA+$U$ by varying the value of $U$ between 0 and 6 eV. We also show the corresponding LCW back-folded results in Figure 5. One can clearly see the similarity between LaB$_6$ and CeB$_6$, $U = 4$ and 6 eV spectra. This can be attributed to the on-site Coulomb repulsion (Hubbard $U$), which pushes unoccupied 4$f$ manifold above the Fermi level and localizes the remaining filled 4$f$-orbitals. The localization of a single electron in the 4$f$-orbital causes the fully ferromagnetic ground state with the magnetic moment of about 1 $\mu_B$.

The shown spectra agree with the 2D-ACAR measurements presented in the study by Biasini et al.\cite{19} for both structures (LaB$_6$ and CeB$_6$). We observe that the CeB$_6$ spectra are similar to LaB$_6$ for $U = 4$ and 6 eV in all shown high-symmetry projections. This effect is connected to the localization of the 4$f$-orbital as its position in energy is shifted further below the FS with increasing values of the Hubbard $U$ parameter. On the central panel of the Figure 5 (001 projection), where we show the LCW back-folded unconvolved data, one can clearly identify...
the FS “ellipsoids” in [100] and [010] directions as well as rounded-square cross sections of the same ellipsoids along the [001] direction. This demonstrates that the high-resolution 2D-ACAR spectra can be used to deduce bulk FS features of the $R$B$_6$ family of compounds, which also includes $4f$-valence electron bands.

4. Summary and Outlook

The 2D-ACAR experiments offer answers to some fundamental questions of whether the electron momentum density in a specific material possesses FS breaks. Through this technique one may provide guidance in searching for a suitable theoretical description of the electronic structure and thus help in developing an understanding of the mechanism responsible for the occurrence of specific effects, among them also topological features. Alternatively, the accuracy of any specific electronic structure model can in principle be assessed by comparing the computed 2D-ACAR spectra with the corresponding measurements. On the theoretical side, such comparisons have been already made using the most advanced LDA+DMFT method for simple transition metals.[26]

However, its extension to correlated multiorbital $4f$-systems with strong spin–orbit coupling is not currently available.

In the present study, we carried out 2D-ACAR measurements on single-crystalline LaB$_6$ along three high-symmetry directions. The experimental spectra in both $p$- and $k$-space were compared with LDA calculations and showed good agreement. In particular the quantitative results extracted from the experiment agree excellently with the corresponding theoretical values of the FS along $X$–$M$ and $X$–$R$ directions; the higher experimental value found for $X$–$I$, however, is attributed to the fact that this direction is experimentally not directly accessible.

For another member of the rare-earth hexaboride class, the CeB$_6$ compound with $4f$-states in the valence band, in absence of the recent experimental measurements, the LDA + $U$ method has been applied with different values of the Hubbard parameter $U$. The present study was motivated following the proposal that SmB$_6$, another member of the hexaboride family, may be an interaction-driven topological insulator.[7,43] The evidence for a metallic surface state in this material is largely accepted, and at the same time low-temperature transport[44,45] and (spin-polarized) angle-resolved photoemission spectroscopy measurements[43] point toward a bulk small-gap insulating state. Quantum oscillation measurements[38] found dHvA frequencies, cyclotron masses, and amplitudes that are quite similar to other metallic hexaborides such as LaB$_6$ and CeB$_6$.[46-48] These two compounds exhibit a metallic ground state, involving predominantly conduction electrons with low residual resistivity, and are characterized according to the DFT (LDA) calculations by a multiple-connected FS of distorted ellipsoids. Band structure calculations for these metallic compounds showed that these types of “ellipsoids” are universal FS features for these materials. In addition, the same types of calculations revealed similar features for SmB$_6$, when the Fermi level is shifted by hand from the computed position, which is in the gap, either up in the conduction bands or down in the valence bands.[38]

Based on the presented results and the combined theoretical and experimental (for LaB$_6$) analysis, we conclude that the “ellipsoid” cross sections ($\sigma$-orbit) and neck sizes increase; hence, $\gamma$- and $\epsilon$-hole orbits of LaB$_6$ and CeB$_6$ are significantly reduced in size in comparison with SmB$_6$. This is in agreement with the quantum oscillation dHvA measurements and supports the idea that these two compounds are topologically trivial but correlated metals. Therefore, as a complementary approach to the present LDA($+U$) treatment of LaB$_6$ and CeB$_6$, we might consider the DMFT which is able to take into account the full local correlation effects and its extensions also including intersite correlations. Here the role of the crystal electric field becomes apparent. The sixfold degenerate $j = 5/2$ multiplet state of Ce-$4f^3$ is split into a $\Gamma_8$ quartet and the $\Gamma_7$ doublet. An effective Hamiltonian formulation restricted to the $\Gamma_8$ quartet only represents a significant reduction of the local Hilbert space and therefore makes the many-body DMFT computation feasible. This will finally allow the description of a truly paramagnetic metallic ground state and may reveal more subtle effects connected to the shape of the FS.

Figure 5. Unconvolved LCW folded-back $k$-space density of the calculated ACAR spectra for LaB$_6$ (upper left) and CeB$_6$ for $U = 0$ (upper right), $U = 4$ eV (bottom left), and $U = 6$ eV (bottom right). Results correspond to LDA (LaB$_6$ and CeB$_6$, $U = 0$) and LDA + $U$ (the rest) calculations with the Drummond parametrization of the electron–positron enhancement factor. The borders of the projected first BZ (white lines) in the repeated zone scheme are shown in all plots. From left to right, (110), (001), and (111) high-symmetry projections.
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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

The data that support the findings of this study are available from the corresponding author upon reasonable request.

Keywords

electronic structure, positron annihilation, rare-earth hexaborides

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