Yb Valence Change in Ce$_{1-x}$Yb$_x$CoIn$_5$ from spectroscopy and bulk properties

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The electronic structure of Ce$_{1-x}$Yb$_x$CoIn$_5$ has been studied by a combination of photoemission, x-ray absorption and bulk property measurements. Previous findings of a Ce valence near 3+ for all x and of an Yb valence near 2.3+ for x ≥ 0.3 were confirmed. One new result of this study is that the Yb valence for x ≤ 0.2 increases rapidly with decreasing x from 2.3 toward 3+, which correlates well with de Haas van Alphen results showing a change of Fermi surface around x = 0.2. Another new result is the direct observation by angle resolved photoemission Fermi surface maps of ≈ 50% cross sectional area reductions of the α and β sheets for x = 1 compared to x = 0, and a smaller, essentially proportionate, size change of the α sheet for x = 0.2. These changes are found to be in good general agreement with expectations from simple electron counting. The implications of these results for the unusual robustness of superconductivity and Kondo coherence with increasing x in this alloy system are discussed.

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I. INTRODUCTION

A. General Overview

Heavy fermion (HF) systems are characterized by a delicate interplay of localized and itinerant electronic degrees of freedom that is responsible for a myriad of interesting strongly correlated electron phenomena. Here the localized f-electrons of Ce, Pr, Yb and various 5f elements embedded in an intermetallic compound hybridize with the conduction electrons.

The physics of the dilute limit of a single f-electron impurity in a metallic host is well-understood and described by the single-ion Kondo problem, where below the single ion Kondo temperature $T_K$, the spins of the conduction electrons quench the local magnetic moment of the impurity via the Kondo interaction. HF compounds represent the dense limit where the f-electron elements are arranged on a lattice, and in turn their local magnetic moments are mutually coupled through the conduction electrons by means of the Ruderman-Kittel-Kasuya-Yosida (RKKY) interaction. The competition between RKKY and Kondo interactions is often summarized in a generic phase diagram for HF materials. On one end the RKKY interaction leads to the development of long-range antiferromagnetic (AFM) order. On the other end the Kondo interaction drives the demagnetization of the f-electron state, resulting in a paramagnetic HF state in which the entire lattice of f-electron moments collectively undergoes the Kondo effect and the f-electrons are delocalized into the conduction band. The competing interactions can frequently be tuned by a nonthermal control parameter such as chemical composition $x$, pressure $P$ or magnetic fields $H$. A magnetic quantum critical point (QCP) is observed when the magnetic critical temperature is suppressed to zero.

Magnetic QCPs in HF compounds have continuously attracted scientific interest because in their vicinity the Fermi-liquid paradigm is observed to break down frequently accompanied by the emergence of unconventional superconductivity (SC). Both phenomena originate from the abundance of soft magnetic quantum fluctuations at the QCP, which in the latter case are believed to provide the pairing “glue” for the SC.

In particular, the family of tetragonal “115” systems has been investigated in great detail in order to disentangle the complex interplay between the heavy fermion state, unconventional SC and quantum criticality. The most prominent member of this class is CeCoIn$_5$. CeCoIn$_5$ evinces superconductivity below a critical temperature $T_c = 2.3$ K, and is thought to be situated on the brink of an AFM QCP. This is reflected in the $T$-linear low-temperature electrical resistivity that indicates the presence of AFM quantum critical fluctuations. Here magnetically mediated SC is supported by the observation of a strong spin-resonance in neutron scattering experiments.

Apart from the SCing state, also the heavy normal state and non-Fermi liquid (NFL) properties of CeCoIn$_5$ have been studied extensively by tuning the system via rare earth substitution on the Ce site. Notably, an analysis of transport data for Ce$_{1-x}$La$_x$CoIn$_5$ introduced the notion of a coherence temperature $T^* ≈ 45$ K below which f-electron delocalization is supposed to proceed. Further, for Ce$_{1-x}$R$_x$CoIn$_5$ it was found that both Cooper pair breaking and Kondo-lattice coherence are uniformly influenced by magnetic and nonmagnetic...
rare earth (R) substituents. In contrast, the NFL behavior is strongly dependent on the f-electron configuration of the R ion. A more recent study suggests that the introduction of small amounts of non-magnetic impurities on the Ce (Y, La, Yb, Th) and In (Hg, Sn) site generates an inhomogeneous electronic state, in which the periodicity of the Kondo lattice is disrupted by the impurities. This additionally results in a rapid local suppression of unconventional superconductivity.

These prior results can be contrasted with the properties of a new alloy series Ce$_{1-x}$Yb$_x$CoIn$_5$ for 0 ≤ x ≤ 1 that may provide a fresh view on both the normal and the SCing state in HF compounds. Notably, it was found that the coherence temperature $T^*$, identified via the low-temperature electrical resistivity maximum, is essentially constant over the entire substitution range. This is surprising by itself, but taking into account that the single ion Kondo temperatures $T_K$ for CeCoIn$_5$ and YbCoIn$_5$ differ (see, e.g., Ref. [16]), it also contradicts a recent study that suggests that $T^*$ and $T_K$ generally scale with each other. The apparent stability of the electronic state is also reflected in the lattice parameters that remain nearly constant for x ≤ 0.775, after which phase separation into Yb rich and deficient phases of Ce$_{1-x}$Yb$_x$CoIn$_5$ occurs. Also the magnetic susceptibility is almost unaffected by the substitution of Ce with Yb. The SCing critical temperature $T_c$ decreases linearly with x towards 0 K as x → 1, in contrast with other HF superconductors where $T_c$ scales with $T^*$ (see, e.g., Ref. [17]). Only the low-temperature NFL behavior derived from the electrical resistivity, specific heat and magnetic susceptibility varies with x, even though there is no readily identifiable quantum critical point.

Two different hypotheses have been put forward to explain the remarkable behavior of Ce$_{1-x}$Yb$_x$CoIn$_5$. Based on the robustness of $T_c$ and $T^*$, and most of all the agreement of the observed NFL behavior with the presence of critical valence fluctuations, Shu et al. have proposed a correlated electron state having cooperative valence fluctuations of Yb and Ce. On the other hand, Booth et al. suggested from EXAFS measurements that below the Yb concentration where macroscopic phase separation takes place there is nonetheless a high degree of inhomogeneity in the form of large coexisting interlaced networks of CeCoIn$_5$ and YbCoIn$_5$. It was argued that the YbCoIn$_5$ network would locally influence the physical properties of CeCoIn$_5$, causing the slow suppression of $T_c$. But it seems that one could equally well imagine that the consequence of such large networks could be an unchanged value of $T_c$ because the CeCoIn$_5$ and YbCoIn$_5$ networks would only influence each other at their respective surfaces. In that case, one would expect the $T_c$ in the CeCoIn$_5$ network remains constant as function of Yb concentration, while the superconducting volume fraction for the entire sample would decrease. We also note that for the samples studied by Booth et al. a change in the distances of nearest neighbor ions has been observed for $x > 0.4$ using EXAFS. This may be interpreted as a local precursor of the macroscopic phase separation that has been observed by Shu et al. at $x ≈ 0.8$.

The implications of recent thin film studies are unclear for the issue of homogeneity. It was found for epitaxial superlattices of CeCoIn$_5$/YbCoIn$_5$ that $T_c$ is suppressed by $x < 0.2$, essentially like the behavior for other (R) substituents. For thin films of Ce$_{1-x}$Yb$_x$CoIn$_5$ $T_c$ is suppressed by $x > 0.4$, which is, on one hand, smaller than for bulk crystals, but on the other hand, still larger than for other (R) substituents. One possible interpretation could be that the films are more homogeneous than the bulk crystals and thus show $T_c$ suppression more quickly. But the more likely possibility is that the thin films constitute an essentially different materials system from the bulk crystals, e.g., owing to the effect of the interaction of the film with the substrate. For the thin films of Ce$_{1-x}$Yb$_x$CoIn$_5$ it was found that the in-plane lattice parameter is expanded slightly because it is in registry with the substrate and does not change with x, whereas the out-of-plane lattice parameter varies linearly with x between the values for CeCoIn$_5$ and YbCoIn$_5$, quite different from the behavior of the bulk crystals. The comparison of thin films to bulk crystals is further complicated by the recent finding that thin films are extremely sensitive to air and degrade quickly upon exposure.

Both hypotheses, the cooperative valence fluctuations as well as the coexisting interlaced networks, have interesting implications and justify a more detailed microscopic investigation of the electronic structure, and in particular the Ce and Yb valences. The possibility of critical valence fluctuations within an extended SCing phase is remarkable in the context of recent studies in which CeRhIn$_5$ and CeCu$_2$Si$_2$ were suggested as candidates for valence-fluctuation mediated SC. Further a recent study of the transport properties of CeRhIn$_5$ under hydrostatic pressure found that scattering of the charge carriers near the AFM QCP is isotropic, in contrast to expectations for a classical AFM QCP. This finding was interpreted as a signature of coexisting critical degrees of freedom in both spin and charge channels that could be a source of SC pairing. On the other hand SC in interlaced networks of CeCoIn$_5$ and YbCoIn$_5$ is of interest in view of a current proposal that unconventional SCs in the vicinity of AFM may be generally electronically textured. Finally, the $x$-dependence of $T^*$ has been addressed in a recent theoretical work which demonstrates that the onset of coherence is strongly affected by the degree of correlations between impurity sites.

### B. Issues of valence and electron counting

With the proposal of cooperative valence fluctuations, the $x$-dependencies of the Ce and Yb valences are of immediate interest. From spectroscopic information and analysis of bulk properties the $x = 0$ compound is known to have essentially trivalent Ce ($4f^1$) and the uncor-
lated behavior of the $x = 1$ compound might suggest divalent Yb ($4f^{14}$). But the picture of cooperative valence fluctuations in the alloy would require intermediate valence Yb. This picture has been supported by a report from Booth et al. from X-ray absorption spectroscopy (XAS) at the Yb and Ce L$_{11}$ edges that the Yb and Ce valences are 2.3 and 3.1, respectively. Ref. [10] also finds these valences to be essentially independent of $x$. The $x = 1$ compound can then be interpreted as intermediate valence with the $4f^{14}$ magnetic moment quenched on such a high energy scale $T_K$ that Curie Weiss behavior cannot be seen. $T_K$ was estimated in Ref. [10] to be larger than 6000 K.

There are also important questions of electron counting and the implications for the volume contained by the Fermi surface (FS). It must be kept in mind that if a local moment is quenched, e.g., by the Kondo effect, then the electrons producing the moment must be included in the Fermi surface (FS). Thus for Ce$^{3+}$ with its magnetic moment Kondo quenched, the FS volume is based on an atomic configuration $[\text{Xe}]4f^{15}5d^16s^2$ having 4 delocalized electrons/Ce, the same as if Ce were formally Ce$^{3+}$ $[\text{Xe}]4f^{9}5d^56s^2$. For CeCoIn$_5$ de Haas van Alphen (dHvA) measurements$^{25-27}$ at low temperatures have found that the 4f electron is included in the Fermi surface although angle resolved PES (ARPES) performed at 20K has found the FS expected in LDA band calculations with the Ce 4f electron confined to the core, i.e. $[\text{Xe}]4f^{15}5d^16s^2$ with 3 electrons/Ce going into the FS$^{25-26}$ Analogously for Yb$^{3+}$ with $4f^{13}$, if the magnetic moment of the 4f hole is quenched, the Fermi surface must contain the 4f hole and so its volume will be the same as though Yb were formally divalent $[\text{Xe}]4f^{14}6s^2$, i.e., the FS volume would contain 2 electrons/Yb. Because of its large $T_K$ this situation is expected up to very high temperatures for YbCoIn$_5$. Thus, no matter whether or not the 4f electron in CeCoIn$_5$ is localized at the measurement temperature, one expects that in comparing the $x = 1$ and the $x = 0$ compounds, characteristic hole FS features will tend to expand and characteristic electron FS features will tend to shrink with increasing $x$.

Recent dHvA experiments$^{30}$ found that the FS of YbCoIn$_5$ is indeed much different from that of CeCoIn$_5$. The measured frequencies are in reasonable agreement with an LDA calculation for YbCoIn$_5$, which gives an Yb valence of 2.3, the same as found in XAS$^{29}$ and shows reduced volume relative to that of $x = 0$. For example the frequencies assigned to two prominent electron FS features centered on the M-A line in $k$-space and known as $\alpha$ and $\beta$ from LDA$^{25,27}$ and dHvA$^{25,27}$ studies for $x = 0$, decrease markedly from $x = 0$ to $x = 1$. It is also of note that the dHvA effective masses for YbCoIn$_5$ are relatively small, in the range 1.0 $m_e$ to 1.5 $m_e$. These small masses are consistent with the very large value of $T_K$ implied by the Yb valence of 2.3. For $x = 0.1$, the dHvA frequencies and masses are unchanged from those of $x = 0$ and for $x = 0.2$ there appear frequencies characteristic of both $x = 0$ and $x = 1$. For $x = 0.55$, the next highest value for which dHvA data were obtained, and for higher values $x = 0.85$ and 0.95, the frequencies and masses that could be observed are generally like those that are found for $x = 1$, with the $\alpha$ frequencies essentially unchanged and the $\beta$ frequencies changing slightly. Thus dHvA shows a rather abrupt change of electronic structure around $x = 0.2$. In contrast, Ref. [16] reported from ARPES that the electronic structure along the Γ-M line is essentially invariant with $x$, including $x = 1$.

C. Present work and Organization of the Paper

In the present work we determine the Ce valence from XAS at the Ce M$_{3,5}$ edges and the Yb valence from 4f electron x-ray photoemission spectroscopy (XPS). In agreement with Ref. [16] we find that the Ce valence is near 3+ and essentially independent of $x$. The Yb valence for $x = 1$ and decreasing to 0.3 is 2.3, also in agreement with the finding of Ref. [16]. But as $x$ decreases further and approaches 0 the valence increases to nearly 3+. We also report and analyze the $x$-dependence of the alloy magnetic susceptibility for temperatures where it exhibits Curie-Weiss behavior, and we introduce a simple model to inter-relate the effective moment and the Ce and Yb valences. Under the assumptions that Kondo effects can be ignored and that the Ce valence is essentially 3+, we infer Yb valences in very good agreement with the values found from XPS, including their tendency to increase toward 3+ for small $x$. Thus the rather abrupt change of electronic structure found in dHvA can now be seen as resulting from a change of Yb valence.

We present the $k$-dependent electronic structure and FS of CeCoIn$_5$ and YbCoIn$_5$ throughout the Brillouin zone as measured using variable photon energy ARPES. In contrast to the results of Ref. [16] we find along the Γ-M line a large difference of electronic structure for $x = 0$ and $x = 1$. In particular the sizes of the $\alpha$ and $\beta$ sheets decrease markedly from $x = 0$ to $x = 1$, in good qualitative agreement with the general expectations from electron counting set forth above and with the dHvA results. But in disagreement with dHvA is the observation of a smaller, essentially proportionate, size change of the $\alpha$ sheet for $x = 0.2$. The somewhat columnar shapes of these FS features have drawn attention in connection with the idea that the layered crystal structure may be important for the SC of the $x = 0$ compound.

The remainder of the paper is organized as follows. The experimental details on the sample preparation, bulk property measurements and spectroscopic measurements are summarized in section IV. Section V presents the various spectroscopic studies and the data analysis used to determine the Yb and Ce valences. In section VI we relate the Ce and Yb valences to the dependence of the magnetic susceptibility and the unit cell volume on the Yb concentration $x$. Our ARPES results for $x = 0$, 0.2 and 1 are presented in section VII. We end with a
II. EXPERIMENTAL DETAILS

Single crystals of Ce$_{1-x}$Yb$_x$CoIn$_5$ were grown using an indium self flux method. High purity elements (Ce, 3N; Yb, 3N; Co, 3N; In, 4N) were placed in aluminia crucibles and heated in quartz tubes with 150 psi argon gas. The heating schedule consisted of an initial ramp at 50 °C/hr to 1050 °C, a dwell at 1050 °C for 72 hours, and a two-stage cooling process to avoid forming crystals of CeIn$_3$ - first a rapid cooling from 1050 °C to 800 °C followed by a slow cool to 450 °C, where the excess flux was spun off in a centrifuge. The resulting crystals were characterized with x-ray powder diffraction (XRD) and energy dispersive x-ray (EDX) analysis in order to verify both the correct structure and composition. The magnetization $M_{ab}$ of the crystals was measured as a function of temperature for $2 \leq T \leq 300$ K using a Quantum Design superconducting quantum interference device (SQUID) magnetometer in a magnetic field $H = 5000$ Oe applied parallel and perpendicular to the basal tetragonal plane.

The spectroscopic measurements XAS, XPS and ARPES were performed at undulator beamline 7.0.1 of the Advanced Light Source (ALS) synchrotron. XAS was measured using total electron yield (TEY), by simultaneously measuring the sample current and the reference of a Ni mesh located before the sample in the monochromator. A Scienta R4000 electron spectrometer with 2D parallel detection of electron kinetic energy and angle in combination with a highly-automated six-axis helium cryostat goniometer was used to acquire Fermi surface (FS) and electronic structure maps with a wide >30 ° angular window covering multiple Brillouin zones (BZs). The measurements were performed with pressure between 8x10$^{-11}$ mbar and 1x10$^{-10}$ mbar. The samples were cleaved in situ by pushing against a post which was glued on the sample surface by epoxy adhesive. The cleavage temperature was between 20 K and 25 K, essentially equal to the measurement temperature, which was 25 K for CeCoIn$_5$ and 20K for YbCoIn$_5$. The position of the Fermi energy and the energy resolution were determined by measuring a gold foil adjacent to and in good thermo-electrical contact with the sample. Before measuring, the gold foil was scraped in situ to obtain a clean surface. Within about 100 μm all spectroscopic data were collected on the same spot of the cleavage plane, which showed no visible inhomogeneities either in situ or afterwards in images in both an optical microscope and an electron microscope.

XAS measurements had a resolution of 250 meV. For XPS, the overall energy resolution was set to 140 meV FWHM for photon energies of $h\nu = 550$ eV and 250 meV for photon energies of $h\nu = 865$ eV. For ARPES the total energy resolution of the analyzer and exciting photons varied from 30 meV at $h\nu = 80$ eV to 45 meV at $h\nu = 200$ eV, and the angular resolution of 0.3° corresponds to a parallel angular momentum resolution range of 0.024 Å$^{-1}$ to 0.037 Å$^{-1}$. Detector angular distortions are corrected using calibration data acquired with a slit array placed between the sample and analyzer lens. Angular and photon-dependent Fermi-energy maps were extracted with an energy width of 50 meV. The value of k perpendicular to the sample surface ($k_z$) could be selected by varying the photon energy, as verified and calibrated from repeating features in $k_z$-$k_x$ maps using a standard method that approximates the photoelectron dispersion by a free electron parabola and an “inner potential” $V_0$ to characterize the surface potential discontinuity. A $V_0$ value of 11.9 ± 0.6 eV best describes the repeating features in the data.

The actual compositions $x_{act}$ of the Ce$_{1-x}$Yb$_x$CoIn$_5$ samples were determined by X-ray energy dispersive spectrometry (XEDS) analysis. Although $x_{act}$ can vary...
considerably from the nominal composition $x_{\text{nom}}$, as described further below, nonetheless it is possible to find samples in which the two are similar and such samples were carefully selected for obtaining the susceptibility data reported here and in Ref. 14. The XEDS analysis for these samples was performed at the University of California at San Diego. The XEDS analysis of the samples used for the synchrotron electron spectroscopy was performed at the Electron Microbeam Analysis Laboratory at the University of Michigan. Since the synchrotron spectroscopy is performed with a small photon spot on a selected region of a cleaved surface the XEDS analysis was performed after the synchrotron spectroscopy was done and great care was taken to obtain the composition at the same location on the sample as that where the synchrotron spectroscopy was performed. Using the focused probe of the scanning electron microscope it is possible to determine the composition of the sample with a resolution of a few μm (the actual value depends on the average atomic number of the sample and the accelerating voltage of the microscope (30kV in this case)). Therefore by this method we can detect inhomogeneities on the micro-scale but not the nano-scale range. As the photon beam-spot for the synchrotron spectroscopy was much larger, roughly 50-100 μm, we checked the micro-scale homogeneity by doing XEDS analysis around the measured position, at two or three points roughly within a circle of 100-200 microns diameter. We detected no significant changes in the Yb compositions, determined as we describe in detail next.

Fig. 1 (a) and (b) illustrate the procedure for a sample with $x_{\text{nom}} = 0.4$. In the synchrotron experiment we obtain the real space sample surface “XY” map displayed in (a) by measuring the photoelectron intensity in a window of 200 meV around the Fermi-energy while scanning the sample position in the X- and Y-directions perpendicular to the sample-analyzer axis. Brighter color means higher photoelectron intensity. We then compare the XY-map with the scanning electron microscope image of the sample and identify the position where we performed the electron spectroscopy. This enables us to perform the XEDS analysis at very much the same position. After obtaining the X-ray fluorescence spectrum, the composition analysis was performed by using the intensity of the In L-lines, the Ce L-lines, the Co K-lines and the Yb L lines. The first step is to perform a so-called “standard-free” analysis that determines the intensities of the multiple lines of the spectrum and then applies a correction in order to account for the deviations of the spectrum of a pure element relative to that of the element residing in the matrix of other elements such as in Ce$_{1-x}$Yb$_x$CoIn$_5$. We used the so-called ZAF method. This method includes an atomic number correction (Z) which estimates the backscattering and stopping power of the incident electron beam, an absorption correction (A) which corrects for absorption within the matrix, and a correction for fluorescence (F) within the matrix. The corrected spectroscopic intensities are then used to obtain relative elemental compositions subject to the constraint that there are 7 atoms per formula unit. For In and Co compositions so obtained were typically near to the expected values of 5 and 1, respectively, but with some outlier values that for In were at most 2.6% different from 5. The magnitude of the maximum outlier discrepancy for the Co concentration was somewhat less than that for the In composition, resulting in a maximum percentage deviation of 9%. We take these discrepancies as an indication of the uncertainties inherent in the technique. The standard free Ce and Yb compositions so obtained for the Ce $(y_{sf})$ and Yb $(x_{sf})$ concentrations of samples with various $x_{\text{nom}}$ values are listed in Table I. In the next step the end-members CeCoIn$_5$ and YbCoIn$_5$ were used as standards in order to account for possible systematic errors in the ZAF procedure, i.e. we would expect our true $y=1$ in CeCoIn$_5$ and our true $x=1$ in YbCoIn$_5$. While the Ce component requires no correction (as CeCoIn$_5$ gives $y_{sf}=1$), the discrepancy for YbCoIn$_5$ results in the correction formula $x_{\text{act}} = x_{sf}/0.92$. For simplicity in the rest of the paper we use $x$ to mean the actual concentration $x_{\text{act}}$ so determined. We show error bars for $x$ estimated as $\Delta x_{\text{act}} = |x_{\text{act}} + y_{sf} - 1|$, reflecting the assumption that all Yb-atoms substitute for Ce-atoms and, therefore, the determination of $(1-y_{sf})$ should be also an equally valid determination of the Yb content. In Fig. 1 (c), we plot the nominal Yb-concentration vs. the actual Yb-concentration. One can see that, beside $x_{\text{nom}} = 1$, all samples in our spectroscopic study are below the phase separation region which is between $x_{\text{nom}} = 0.775$ and $x_{\text{nom}} = 1$. We compared our results (circles) with the results of Capan et al, who performed XEDS on single crystals and also proton-induced X-ray emission microprobe (PIXE) on a mosaic of crystals from the same batches. One sees clearly a good agreement between our results and the ones of Ref. 14 except for two outlier $x_{\text{nom}}$ values of the latter data. This agreement with independent results from two other techniques encourages us to have confidence in our procedure. We note that the general conclusions drawn in our paper are made with full cognizance of our error bars and do not depend on whether or not the systematic Yb correction described above was made.

| Element | Nominal composition $x_{\text{nom}}$ |
|---------|----------------------------------|
| Ce content $y_{sf}$ | 1.00 | 1.01 | 1.01 | 0.95 | 0.77 | 0.87 | 0.00 |
| Yb content $x_{sf}$ | 0.00 | 0.04 | 0.04 | 0.11 | 0.24 | 0.19 | 0.92 |
| $x_{\text{act}}$ | 0.00 | 0.04 | 0.04 | 0.12 | 0.26 | 0.21 | 1.00 |
| $\Delta x_{\text{act}}$ | 0.00 | 0.05 | 0.05 | 0.05 | 0.03 | 0.08 | 0.00 |

TABLE I: Results of the XEDS analysis.
III. CE AND YB VALENCES FROM SPECTROSCOPY

A. XAS for Ce valence

In order to determine the change of the Ce valence upon doping, we performed TEY XAS near the Ce M\(_4\) and M\(_5\) edges. In this experiment we look at the 3d\(^{10}\)f\(^0\) \(\rightarrow\) 3d\(^7\)f\(^3\) and 3d\(^{10}\)f\(^1\) \(\rightarrow\) 3d\(^7\)f\(^2\) absorption lines, which are further separated accordingly to the core hole being 3d\(_{5/2}\) (M\(_5\)) or 3d\(_{3/2}\) (M\(_4\)). Because the d core-hole interacts strongly with the promoted f-electron, this absorption results in strong excitonic lines at energies below the true 3d \(\rightarrow\) 6p absorption edges, with characteristic structure due to multiplet splittings of the final states. In the two topmost curves of Fig. 2(a), we show an atomic multiplet calculation\(^{39}\) for the initial state (final state) being f\(^0\) (f\(^1\)) or f\(^1\) (f\(^0\)).

A mixed valence system shows absorption lines of both valences. The intensity ratio \(R = I(f^0)/[I(f^0) + I(f^1)]\), where I(f\(^0\)) and I(f\(^1\)) are the integrated intensities for lines with initial states f\(^0\) and f\(^1\), respectively, is a measure of the initial state f\(^0\) component (1-\(n_{f}^{Ce}\)), where \(n_{f}^{Ce}\) is the Ce 4f occupancy. There are, however, the following caveats. First, within the framework of the Anderson impurity model, \(R\) underestimates (1-\(n_{f}^{Ce}\)) due to the mixing of the f\(^1\) and f\(^2\) final states through the hybridization of the f-states with the conduction electron states\(^{37,38}\). Second, TEY detection has a contribution from the surface, which can have smaller (1-\(n_{f}^{Ce}\)) than the bulk because the hybridization can be reduced and the 4f binding energy increased relative to the bulk. Generally, the total fluorescence yield is more bulk sensitive than TEY and also tends to enhance the f\(^0\) peak due to the bulk self-absorption of the much stronger Ce f\(^1\)-component. The greatest sensitivity to the f\(^0\) component is achieved through XAS on L\(_{III}\) edges and resonant x-ray emission\(^{39}\). Thus really precise absolute values for the Ce valence are not accessible here. Nonetheless the M\(_{4,5}\) spectra are known to be a very reliable qualitative guide to (1-\(n_{f}^{Ce}\)) and are very accurate for the main purpose here of detecting a relative change with \(x\) if it exists.

The lower curves in Fig. 2(a) show experimental XAS results, all at \(T = 20\) K, for the series Ce\(_{1-x}\)Yb\(_x\)CoIn\(_5\). The curve for \(x = 0\) is from Ref. \(^{10}\) and indicates therefore reproducibility of the results. Our spectra were normalized at the background from \(h\nu = 910\) eV to \(h\nu = 915\) eV. The spectrum for \(x = 1\) shows expectedly no Ce signal. For \(x < 1\), the spectra are dominated by the f\(^1\)-component with only a small f\(^0\)-satellite. In Fig. 2(b), we show a magnification of the low intensity region. At about \(h\nu = 888\) eV and \(h\nu = 906\) eV, there are little lumps due to the f\(^0\)-component. Also one can see in that magnification that the (dotted) curve from the atomic multiplet calculation for f\(^0\) has to be shifted by about +2.5 eV to account for increased screening in the solid state. Overall, the measured curves do not strongly change with \(x\). We can safely conclude, in agreement with Ref. \(^{10}\) that Ce is essentially trivalent andunchanging for the whole measured series. As a quantitative measure of a possible valence change of the Ce, we determined the intensity of the f\(^1\)-component by a fit with five Gaussians for M\(_5\) and four Gaussians for M\(_4\). Similarly, we determined the f\(^0\)-component by one Gaussian each for M\(_5\) and M\(_4\). Thereby we find at \(T = 20\) K that the value of \(R\) is essentially constant at 0.04 ± 0.04 and 0.1 ± 0.12 for M\(_5\) and M\(_4\), respectively.

B. 4f PES for Yb valence

In order to elucidate the valence of the Yb, we analyze 4f XPS spectra. At the beginning of our measurements, we routinely confirm the absence of oxygenated surfaces by taking scans over a large binding energy region that includes the core-states as shown in the inset.
of Fig. 3 (a). These wide scans also offer the opportunity to normalize our valence-band spectra by assuming that each photon energy the area of the In-3d peaks is constant for the whole series of Ce$_{1-x}$Yb$_x$CoIn$_5$. We show such normalized valence band spectra for the photon energy of $h\nu=865$ eV in Fig. 3 (a). The data are stacked with a constant shift in intensity and are ordered with increasing Yb concentration from bottom to top, i.e. the lowest is $x=0.04$ followed by $x=0.05$, 0.12, 0.21, 0.26, 1. The first notable feature, which we label in the spectra, is the final state $^2F_{5/2}$ - $^2F_{7/2}$ doublet of the Yb$^{2+}$ photoemission process $4f^{14}$ $\rightarrow$ $4f^{13}$. This feature is quite intense because the large number of fourteen Yb $f$-electrons causes a much stronger photoemission signal compared to those of Ce, In, or Co. Tuning the photon energy to the $M_5$ resonance at $h\nu=883$ eV (compare also with Fig. 2) allows us to see the Ce-weight as shown in Fig. 3 (b). As expected we see that this Ce weight decreases upon adding more Yb although the Yb$^{2+}$ doublet is still very strong as seen by comparing the intensity for the $x=1$ sample, which certainly has no Ce weight, with those of Ce, In, or Co. Tuning the photon energy to the $M_5$ resonance at $h\nu=883$ eV (compare also with Fig. 2) allows us to see the Ce-weight as shown in Fig. 3 (b). As expected we see that this Ce weight decreases upon adding more Yb although the Yb$^{2+}$ doublet is still very strong as seen by comparing the intensity for the $x=1$ sample, which certainly has no Ce weight, with the Ce-signal enhanced intensity for all spectra for $x<1$. Going back to Fig. 3 (a), the second notable feature is two lower intensity peaks which are marked by 'S' above them. These two peaks show the same energy separation between each other as the separation of the $^2F_{5/2}$ -
$^2\text{F}_{7/2}$ doublet described above. Thus they are also due to Yb$^{2+}$ but instead of coming from the bulk this signal comes from the surface. The absence of ligand-atoms causes a stronger screening of the $f$-levels and therefore the spectrum shifts down in energy. Such surface shifted Yb$^{2+}$ peaks are well known.

The third feature in the spectrum, finally, is of lower intensity and can be better seen in Fig. 3 (c), which shows a magnification of the interesting region. Although the structure consists of many peaks their mutual origin is from Yb$^{3+}$ for which the photoemission process $5f^{13}\rightarrow 5f^{12}$ gives a final state multiplet of thirteen lines. The topmost curve serves as a fingerprint for Yb$^{3+}$ as it represents a calculation of this multiplet structure using intermediate coupling. Counting the lines of this multiplet, one finds only twelve. The missing line is from a $5f/2^-$ level at 16.3 eV binding energy, which causes it to be merged with the strong signal of the In 4$d$ and Ce $5p$ doublets. For our later quantitative discussion, it is good to note that, according to the calculation, this line has only about 0.09/13 0.7 % of the total intensity of the multiplet and is therefore quite negligible. We notice in Fig. 3 (c) that, as we would expect, the intensity of the Yb$^{3+}$ signal decreases as the concentration of Yb decreases. This however does not necessarily mean that the Yb valence has to change.

Before we start to evaluate the Yb valence qualitatively, we note that there is a relatively easy method to graphically visualize the Yb valence by normalizing the spectrum into components from Yb$^{3+}$ and $^{3+}$Yb. For these extra peaks, we took just enough extra peaks to optimally fit the spectrum. This is even true without taking account of the fact that the normalization overestimates the Yb$^{2+}$ component for small $x$. There is one feature which is more revealed as the absolute intensity of the Yb-related features is more and more decreased. It is marked with an arrow and it stems from the Co $d$-states. This growing of the Co-$d$ related weight shifts the edge "E" of the foremost part of the Yb$^{3+}$ multiplet up. Taking the height between the edge "E" and the dip located between E and the Co-$d$ peak, we can qualitatively state that for $x=1$ the Yb$^{2+}$ component is strong. We can also clearly see that the Yb$^{3+}$ component has a lower intensity for $x=1$ and $x=0.26$ than it has for $x<0.26$. This is even true without taking account of the fact that the normalization overestimates the Yb$^{2+}$ component for low $x$.

In order to quantitatively extract the Yb valence, we applied a fitting procedure presented in Fig. 4. The background used was a Shirley-like background and, additionally, a linear background which simulates the contribution coming from the In 4$d$ and Ce $5p$ doublets at higher binding energies. We modeled the $F_{5/2} - F_{7/2}$ doublet of the bulk Yb$^{2+}$ as two Lorentzians with the fixed intensity ration 6:8. The Yb$^{2+}$ surface component was modeled similarly. We reduced the twelve Yb$^{3+}$ $5f^{13}\rightarrow 5f^{12}$ lines to 7 Lorentzians. The weights and positions of these Lorentzians relative to each other were determined at $x = 1$ (see inset of $x = 1$ in Fig. 4). For fitting the other Yb concentrations, we allow only the relative intensities of the two valence components to change. The fitting routine first finds the bulk Yb$^{2+}$-component and the Yb$^{3+}$ component together with the background, adds then the surface Yb$^{2+}$, and after that subsequently adds just enough extra peaks to optimally fit the spectrum. These extra peaks are mainly to simulate the contributions of Co-$d$ and Ce-$f$. For these extra peaks, we took two or three Lorentzians. All spectral features were con-
volved with a Gaussian having the FWHM of the resolution. Having so many components for a line-fit, we may not always correctly distinguish between the surface Yb$^{2+}$-component and these extra peaks. However, as can be seen in Fig. 4 the peaks labeled as ‘rest’ represent very much what would be expected from the Co and Ce weights. For these Ce weights, the reader can compare with Fig. 3 (c).

The result of the fitting procedure is summarized in Fig. 5. There, we plot the fitting result for both photon energies. The fact that the results are essentially the same for both photon energies assures us that the fit is finding the amount of the surface Yb$^{2+}$-component correctly and that the valence obtained here reflects that of the bulk. For the general trend of the valence vs. Yb concentration we see that the Yb valence is near 3+ at very low $x$ and goes down to about +2.3, as already concluded qualitatively from Fig. 3. Compared to the results of Ref. 16, we obtain the same valence of about +2.3 for the end-member $x = 1$. As discussed in Ref. 16, the intermediate valence of the end-member together with the nonmagnetic behavior seen in its magnetic susceptibility indicates that YbCoIn$_5$ has a very high single ion Kondo temperature $T_K$. However, by measuring for lower $x$-values than in Ref. 16, we obtain the new result that the Yb valence is strongly increasing to trivalent for small values of $x$ going to zero. We will see in the next section that an analysis of the valence from bulk properties is consistent with the rather abrupt increase of valence at low $x$ observed spectroscopically. As noted already in Section IV this change of Yb valence is also in good agreement with the change of electronic structure observed in dHvA experiments.

**IV. RELATION OF CE AND YB VALENCES TO BULK PROPERTIES**

![Graph showing inverse magnetic susceptibility vs. temperature](image)

**FIG. 6:** (a) Inverse magnetic susceptibility $\chi_{ab}^{-1}(T) = H/M_{ab}(T)$ vs. temperature $T$ along the crystallographic $ab$ plane for Ce$_{1-x}$Yb$_x$CoIn$_5$ with Yb concentrations $0.0 \leq x \leq 0.775$. Curie-Weiss temperature $\theta_{CW}$ and effective magnetic moment $\mu_{eff}$ as determined from fits to a Curie-Weiss law $\chi_{ab} = N_A\mu_{eff}^2/3k_B(T - \theta_{CW})$ (please see text) as function of Yb concentration $x$ are shown in (b) and (c), respectively. Solid line in (c): linear fit of $\mu_{eff}(x)$.

Figure 6(a) shows the inverse magnetic susceptibility $\chi_{ab}^{-1}(T) = H/M_{ab}(T)$ of Ce$_{1-x}$Yb$_x$CoIn$_5$ in the normal state vs. Yb concentration $x$. The data were collected by warming the sample gradually after zero-field cooling (ZFC), and subsequently field cooling (FC). The difference between the ZFC and FC data is negligible. Above $T = 30$ K and for all $x$, the magnetic susceptibility $\chi_{ab}$ of Ce$_{1-x}$Yb$_x$CoIn$_5$ can be described well by a Curie-Weiss law

$$\chi_{ab} = N_A\mu_{eff}^2/3k_B(T - \theta_{CW}),$$

where $N_A$ is Avogadro’s number and $k_B$ is Boltzmann’s constant. The effective magnetic moment $\mu_{eff}$ and the Curie-Weiss temperature $\theta_{CW}$ as determined from fits of the data (solid lines in Fig. 6(a)) to Eq. 1 are shown in Fig. 6(b) and (c). The fits yield $\theta_{CW} \approx -120$ K, independent of Yb concentration to first approximation (Fig. 6(b)). Curie-Weiss behavior is also observed in temperature dependence of magnetic susceptibility $\chi_c$ at high temperatures ($T \gtrsim 50$ K), and the fits give similar values of $\mu_{eff}$ and $\theta_{CW}$.

The magnetic susceptibility of metals containing lanthanide ions that exhibit the Kondo effect or valence fluctuations can be described by a Curie-Weiss law (Eq. 1) in the high temperature limit. For the present situation, the effective magnetic moments of the Ce and Yb ions are expected to be close to their Hund’s rules values.
corresponding to their f-electron configurations and the
Curie-Weiss temperature represents a characteristic tem-
perature associated with the Kondo effect for the triva-
lent Ce ions or valence fluctuations for the intermediate
valent Yb ions. The Kondo and valence fluctuation
temperatures are characteristic temperatures where the
material gradually crosses over from a paramagnetic lo-
calized moment regime at high temperatures to a non-
magnetic Pauli-like regime at low temperatures. In our
analysis, the magnetic susceptibility is taken to be a su-
perposition of a Kondo contribution from the Ce ions and
a valence fluctuation contribution from the Yb ions. The
fact that Curie-Weiss temperature theta CW is nearly in-
dependent of Yb concentration indicates that the energy
scale associated with the combined Kondo and valence
fluctuation contributions does not vary with Yb concen-
tration, which is consistent with the stability of the cor-
related electron state in Ce1−xYbxCoIn5 over a large Yb
concentration range.

The magnetic susceptibility $\chi(x)_{ab}$ of Ce1−xYbxCoIn5
is composed of two contributions arising from Ce and Yb
ions, respectively,

$$\chi(x)_{ab} = \chi_{Ce}(1-x) + \chi_{Yb}x. \quad (2)$$

Using the result that $\theta_{CW}$ is independent of $x$ (cf.
Fig. 6(b)), we can write

$$\mu_{eff}^2(x) = \mu_{Ce}^2(x)(1-x) + \mu_{Yb}^2(x)x. \quad (3)$$

Following the conclusion of the previous section and
of Ref. 16 we take the f-electron orbital occupancy for
Ce to remain close to 1 (n_f^{Ce} = 1), i.e., the Ce ions remain
3+ for all $x$. Thus, by assuming that $\mu_{Ce}(x) = \mu_{Ce3+} =
2.54 \mu_B$ for all $x$ and approximating $\mu_{eff}(x)$ by the
linear fit illustrated in Fig. 6(c), we can estimate $\mu_{Yb}(x)$
by using Eq. (3). The f-hole occupancy for Yb is given by

$$n_f^{Yb}(x) = \frac{n_f^{Yb+}(x)}{n_f^{Yb+}},$$

where $n_f^{Yb+} = 4.54 \mu_B$. Accordingly, the
effective valence is then obtained as

$$v_{Yb}(x) = 2 + n_f^{Yb}(x). \quad (4)$$

As shown by the open circles in Fig. 5 the result is in
surprisingly good agreement with the results of the XPS
measurements, considering the simplicity of analysis of
the magnetic susceptibility.

The analysis of the magnetic susceptibility measure-
ments on the Ce1−xYbxCoIn5 crystals used to estimate
the Yb valence relies on the value of the Yb concen-
tration $x$. Most of the magnetic susceptibility measure-
ments were made on samples for which the composition
measured by XEDS was close to the nominal composition
(within about 5%). Coupled with the uncertainty in the
magnetic susceptibility measurements, we roughly esti-
mate that uncertainties in Yb valence are of the order
of 10%. Samples whose compositions were not measured
by XEDS had values of magnetic susceptibility and $T_c$
that changed systematically with nominal composition,
indicating that the actual compositions are close to the
nominal values. This suggests that the XEDS measure-
ments may not be a reliable method of estimating the
bulk Yb concentration in this system for reasons that are
not presently understood. In addition, we again stress
that the assumptions on which the analysis of the mag-
netic susceptibility measurements is based, and that were
used to estimate the Yb valence, are, while reasonable,
not rigorously justified.

There is a well-known correlation between lattice pa-
rameters and valence in f-electron materials. For the
system Ce1−xYbxCoIn5, which has a tetragonal crystal
structure with basal plane and interplane lattice param-
ers $a$ and $c$, respectively, we consider the relationship
between the unit cell volume $V = a^2c$ and the valences
of the Ce and Yb ions. The lattice parameters $a (c)$ for
CeCoIn5 and YbCoIn5 determined from XRD measure-
ments are 4.6012 Å (7.5537 Å) and 4.5590 Å (7.433 Å),
respectively, and the valence of Ce and Yb as obtained by
our photoemission measurements and calculations are 3+ and
2.3+, respectively. This yields unit cell volumes of $V =
159.9 \text{ Å}^3$ for CeCoIn5 and 154.5 \text{ Å}^3 for YbCoIn5.
Assuming that Vegard’s law applies to the unit cell volume
$V$ as a function of $x$ and that the valence of Ce remains
near 3+ for all values of $x$, $V(x)$ can be expressed as

$$V(x) = V_{Ce3+}(1-x) + V_{Yb}x$$
$$= V_{Ce3+}(1-x) + [V_{Yb3+} + V_{Yb2+} - V_{Yb3+}](1 - n_f^{Yb}(x)) \times$$
$$= 159.9 \text{ Å}^3(1-x) + [155.6 \text{ Å}^3 - 3.6 \text{ Å}^3 n_f^{Yb}(x)]x. \quad (5)$$

The values of the unit cell volumes used in this ex-
pression are $V_{Ce3+} = 159.9 \text{ Å}^3$, $V_{Yb3+} = 155.6 \text{ Å}^3$, and
$V_{Yb2+} = 152.0 \text{ Å}^3$, while $n_f^{Yb}(x)$ is the number of holes
in the Yb 4f-electron shell ($n_f^{Yb}(x) = 1$ for Yb3+ and 0
for Yb2+). The value of $V_{Yb3+}$ was estimated by interpo-
larating the values of $V_{Ln3+}$ from the neighboring LnCoIn5
compounds with trivalent Ln ions to YbCoIn5. The
value of $V_{Yb2+}$ was estimated using the value of $V_{Yb3+}$
and $V_{Yb2+}$, inferred from the XAS data reported herein
for the compound YbCoIn5, using the term for $V_{Yb}$ in
brackets in Eq. 5 which yields

$$V_{Yb2+} = [V_{Yb}(x) - V_{Yb3+} n_f^{Yb}(x)] / (1 - n_f^{Yb}(x)). \quad (6)$$
Using \( n_f^{Yb}(1) = 0.3 \) for \( x = 1 \), we obtain \( V_{Yb^{2+}} = 155.6 \, \text{Å}^3 \) from Eq. 4.

In Fig. 7, we compare \( V(x) \) determined from Eq. 5 with the values obtained from the XRD measurements on Ce\(_{1-x}\)Yb\(_x\)CoIn\(_5\). It can be seen that the calculated values of \( V(x) \) are nearly linear and generally conform to the behavior expected from Vegard’s law for Ce\(^{3+}\) and Yb\(^{2.3+}\). The calculated values of \( V(x) \) are smaller than the measured values, and the discrepancy is larger for larger values of \( x \). A number of factors could contribute to this discrepancy; e.g., (1) Weakening of the metallic bond due to the decrease in the conduction electron density with Yb concentration that accompanies the substitution of Yb ions (valence \( 2.3^+ \)) for Ce ions (valence \( 3^+ \)), resulting in an increase of the unit cell volume, (2) a nonlinear contribution considered by Varma and Heine \(^{46}\) in calculating the unit cell volume for Ln compounds with intermediate valence, and (3) a reduction of the actual Yb concentration compared to the nominal concentration.

There are many details visible in these six FS maps and furthermore they represent only a small fraction of the data measured throughout the whole BZ and in multiple zones. Here we will focus only on the \( \alpha \) - and \( \beta \)-sheets which are readily identified in the data. The more complex FS pieces will be analyzed and discussed in a separate publication \(^{47}\). For CeCoIn\(_5\) these complex pieces display topological differences that depend on whether or not \(^{30}\) the FS contains the Ce \( f \)-electron, i.e., whether the FS is "large" or "small." For CeCoIn\(_5\) ARPES finds \(^{23,20}\) that the Ce \( 4f \) electrons behave predominantly localized for the present measurement temperature even though low \( T \) de Haas van Alphen experiments \(^{25,22}\) unambiguously detect the large FS. For the \( \alpha \) - and \( \beta \)-sheets discussed here LDA calculations \(^{50}\) performed for isoelectronic CeRhIn\(_5\) with the Ce \( f \)-electron confined to the core show only small changes in size and no dramatic topological changes due to localizing the \( f \)-electron and excluding it from the FS. Nonetheless we will now see that these sheets display large size changes as \( x \) varies from 0 to 1. For \( x = 0 \) and 1 these size changes are in good agreement with findings from dHvA experiments \(^{30}\) and accompanying LDA calculations for YbCoIn\(_5\). However for the intermediate value of \( x \) there is an important difference from the dHvA results, as discussed below.

The two mesh models at the top right and left in the figure show for \( x = 0 \) the three dimensional shapes of the \( \alpha \)-sheet and the \( \beta \)-sheet, respectively. The mesh models are based on an itinerant LDA calculation and are taken from Ref. \(^{19}\). The \( \alpha \)-sheet is a nearly two-dimensional infinite cylinder along the (001)-direction \((k_z\)-direction\). Measurements using the de Haas van Alphen effect observe three distinct (001) frequencies which LDA identifies as not only the \( \Gamma \)-plane diamond \((\alpha_2)\) and Z-plane square \((\alpha_3)\) orbits but also a slightly larger circular orbit \((\alpha_1)\) just above and below the Z-plane \(^{21}\). Comparing the mesh models with the ARPES measurements, we identify the large round contours centered on the BZ corners as coming from the quasi-two-dimensional \( \alpha \)-sheet. Consistent with this identification, this contour evolves from a diamond-like shape in the \( \Gamma \)-plane of CeCoIn\(_5\) to a slightly squarish shape in the Z-plane. Due to the experimental broadening of ARPES in the \( k_z \)-direction, the observed Z-plane \( \alpha \)-sheet contour has a distinctly broader width with a squarish inside edge and a rounder outside edge, which is interpretable as the contour of \( \alpha_3 \) being blurred with that of \( \alpha_1 \). For YbCoIn\(_5\), we can similarly identify the same shapes as belonging to the \( \alpha \)-sheet. The contours observed in the Z-plane have more of a squarish shape than those of CeCoIn\(_5\), perhaps consistent with LDA calculations \(^{10}\) for YbCoIn\(_5\) that show...
respectively mesh models and 20 K respectively. Top row figures at right and left show the α and β-plane for the Γ and Z planes for FIG. 8: Second, third and fourth rows show Fermi-surface sheets is easily seen. Although the circular shape of the β-sheet is clearly visible by eye in the FS maps is the considerable reduction of size for the α-sheet and the β-sheet. In contrast to the conclusion of Ref. 16 our results show that the electronic structure along the Γ-M line is much different for x = 0 and x = 1, even though the general shapes of the α and β-sheets are much the same. We have made a quantitative analysis to determine the cross-section areas in units of (π/a)², along with the implied dHvA frequencies. The results are listed in Table II along with measured dHvA frequencies for x = 0 and 1. As discussed below a direct comparison to dHvA frequencies for x = 0.2 is not possible. For x = 0 there is good general agreement among the dHvA frequencies for the three studies cited. The ARPES frequencies have a similar pattern of variation in magnitude among the various orbits but are systematically smaller than those from dHvA, a difference for which one can consider two contributing effects. The first is the difference in temperature of the two measurements, that low-T dHvA sees the ’large’ FS and that higher-T ARPES likely sees the ’small’ FS or at least a smaller FS. However dHvA and LDA studies find that the fractional differences occurring in the α and β-sheets for the change from localized to itinerant Ce f electrons in the Ce 115 compounds are relatively small, so this effect may not be very important. Second, even at the ARPES measurement temperature the bands are still heavy enough very near EF over the energy window of the FS maps that the kp value for an electron pocket is likely to be underestimated. For x = 1 neither of these differences are important because the measurement temperatures for dHvA and ARPES are both much less than the characteristic temperature and the bands are light. Indeed here we find much better quantitative agreement between the two techniques. It should however be noted that for the α-sheet the good agreement is aided by assigning the measured dHvA frequencies somewhat differently than in Ref. 39. In that paper the four frequencies labeled F3 through F6 (1.8 kT, 2.04 kT, 2.19 kT and 2.96 kT, respectively) are all associated with the α-sheet but F1 is assigned to α1 and F0 is assigned to α2, leaving F3 and F5 to be assigned to the corregations predicted in LDA between the Γ-plane and the Z-plane. Considering the shape of the LDA α-sheet we find it more natural and in better agreement with ARPES to assign F4 as α1 and F6 as α2, with the other two belonging to the corregations. These reassignments are not inconsistent with the observed angular dependences of the orbits.

Table II also lists values for the ARPES FS areas for x = 1 and x = 0.2 relative to those for x = 0. For the Γ-plane, the ratio between the α-sheet area in YbCoIn5 compared to the area in CeCoIn5 is 0.46, while for the Z-plane this ratio is 0.54. From these data we would estimate that the average ratio of the volumes of the α-sheets is roughly 50%. We note that Ref. 30 also concluded a value of about 50% from the dHvA data, but the combination of the larger dHvA frequencies for x = 0 and our changed orbit assignments for x = 1, as discussed in the preceding paragraph, causes the dHvA data of Table II to
The significant finding for the ARPES data for \( x = 0.2 \) is that the \( \alpha \)-sheet areas are, even by eye, intermediate between those for \( x = 0 \) and \( x = 1 \). Quantitatively the change in area from \( x = 0 \) to \( x = 0.2 \) is 21% and 14% of the change from \( x = 0 \) to \( x = 1 \) for \( \alpha_1 \) and \( \alpha_2 \) respectively, i.e. roughly in the same proportion as the doping. Unfortunately it is not possible to compare this result with the dHvA data for intermediate values of \( x \). Although the frequency \( F_2 \) of the dHvA data is very similar to that implied by the ARPES data (i.e., 3.6 kT or 3.7 kT) this association cannot be made because \( F_2 \) has hardly any variation with \( x \) and in Ref. \[30\] is thought to be associated somehow with the \( \beta \)-sheet. Indeed the dHvA frequencies associated with the \( \alpha \)-sheets and \( \beta \)-sheets of the end members \( x = 0 \) and \( x = 1 \) show only slight modification for intermediate \( x \)-values, quite different from the ARPES finding here of a clear intermediate \( \alpha \)-sheet size for an intermediate \( x \)-value. We have no hypothesis for this very great difference beyond the possibility of some sample difference for intermediate \( x \).

### VI. SUMMARY AND CONCLUSIONS

There are three main new findings of this work for Ce\(_{1-x}\)Yb\(_x\)CoIn\(_5\). First, for small values of \( x \) increasing from 0, the Yb valence changes rapidly from being nearly trivalent to the value of 2.3 found previously for larger \( x \) and for YbCoIn\(_5\). Second, we have directly observed a large reduction in the sizes of the columnar \( \alpha \) and \( \beta \) FS features for \( x = 1 \) relative to those for \( x = 0 \). As already noted, both of these findings are in very good agreement with dHvA results.\[30\] Taken together, these results imply that around \( x = 0.2 \) a change of Yb valence drives a switch of the near \( E_F \) electronic structure from one characteristic of \( x = 0 \) with some very heavy mass FS pieces to one characteristic of \( x = 1 \) with Yb valence around 2.3, a very large Yb \( T_K \) and small measured masses. Third, for at least one intermediate \( x \)-value and one sheet (\( \alpha \)), the FS has been found to evolve between that of the two end members. This third result contrasts quite sharply with the dHvA results, in which the observed frequencies and masses do not evolve significantly with \( x \) and change quickly from one to the other, with a mix of both being seen for \( x = 0.2 \).

What are the implications of these findings for understanding the transport properties? Regarding the two hypotheses mentioned in Section [A] that of cooperative valence fluctuations or coexisting networks of CeCoIn\(_5\) and YbCoIn\(_5\), the ARPES results clearly favor the former, at least for the samples used in this study. The addition of only a small amount of Yb drives an overall change in the FS and near \( E_F \) electronic structure, and there appear to be unified electronic structures presumably involving the f-states of both Ce and Yb. On the other hand the dHvA results lead to a different conclusion. The finding there of aspects of both electronic structures at \( x = 0.2 \) suggests some coexistence. Further it is very puzzling that the observed dHvA frequencies,

| Composition | ARPES | dHvA | Ref. | dHvA | Ref. | dHvA |
|------------|------|------|------|------|------|------|
| Label | FS-Area | FS Area | \( \Delta \nu \) | \( \Delta \nu \) | \( \Delta \nu \) |
| \( x=0 \) | \( \alpha_1 \) | 0.85 ± 0.06 | 1 | 4.1 ± 0.3 | 5.46 (\( \alpha_1 \))/4.37 (\( \alpha_3 \)) |
| | \( \alpha_2 \) | 0.81 ± 0.06 | 1 | 3.9 ± 0.3 | 4.87 (\( \alpha_2 \)) |
| | \( \beta_1 \) | 1.8 ± 0.15 | 1 | 8.7 ± 0.7 | 11.6 (\( \beta_1 \))† |
| | \( \beta_2 \) | 1.12 ± 0.15 | 1 | 5.4 ± 0.7 | 7.4 (\( \beta_2 \))† |
| \( x=0.2 \) | \( \alpha_1 \) | 0.77 ± 0.15 | 0.91 | 3.7 ± 0.7 | |
| | \( \alpha_2 \) | 0.74 ± 0.2 | 0.91 | 3.6 ± 1 | |
| \( x=1 \) | \( \alpha_1 \) | 0.46 ± 0.08 | 0.54 | 2.2 ± 0.4 | 2.2 (\( F_5 \))† |
| | \( \alpha_2 \) | 0.38 ± 0.06 | 0.46 | 1.8 ± 0.3 | 1.6 (\( F_3 \))† |
| | \( \beta_1 \) | 1.1 ± 0.08 | 0.61 | 5.3 ± 0.4 | 6.84 (\( \beta_2 / F_9 \)) |
| | \( \beta_2 \) | 0.5 ± 0.1 | 0.44 | 2.4 ± 0.5 | 3.66 (\( \beta_2 / F_8 \)) |

TABLE II: Left portion: Fermi-surface cross-sectional areas and implied dHvA frequencies from ARPES Fermi-surface maps calculated in units of kilo Tesla (kT); Right portion: Measured dHvA frequencies with original paper labeling shown in parentheses. For \( x = 1 \) Ref. \[30\] reverses the labeling of the \( \beta \)-sheet orbits from that used for \( x = 0 \) such that \( \beta_1 \) corresponds to the Z-plane and \( \beta_2 \) corresponds to the Γ-plane. †Value extracted from a publication graph because not explicitly tabulated.
e.g., for the $\alpha$ orbits, do not change as $x$ increases beyond 0.2, given that the FS size changes appear to be driven by single electron counting and that the total number of electrons to be contained in the FS per rare earth atom certainly changes steadily. Also, as noted in Ref. [30] the switch to a FS with only low mass measured features is inconsistent with the finding that the specific heat is roughly constant with $x$. This issue gave rise to the conclusion[30] that there must be heavy FS pieces for larger $x$ not yet observed in dHvA, possibly the heavier $\beta$ orbits that were not observed for $x = 0.2$ and 0.55. The change in transport properties across the crystallographically two-phase region between $x = 0.8$ and $x = 1$ also bears thought. The absence of Kondo-like features in the resistivity for $x = 1$ is readily understandable from the Yb valence of 2.3, the implied large $T_K$ and the low dHvA masses. But the presence of these features for $x = 0.8$, very similar to those for $x = 0$, plus the lack of any Yb valence change across the region, implies that the resistivity is due only to the Ce and that the change is simply the result of removing all the Ce from the lattice. In this respect the Ce would seem to be acting independently of the Yb.

What is the role of the Ce and Yb for the SC? That $T_c$ decreases only slowly and gradually with $x$ implies a gradual steady change of some essential ingredients for the SC and it is again tempting to think of Ce and Yb as acting independently. One possibility is that Ce brings a local moment which is essential for the SC. In this picture the smallness of the Ce $T_K$ is a benefit, and Yb, with the largeness of its $T_K$ does no direct harm but does serve to dilute the Ce. Another model involving Ce as the essential active ingredient perhaps being diluted by Yb is the composite pairing picture[30]. Moving away from local pictures, if low dimensionality is involved in the SC, as has been suggested for pairing involving spin fluctuations, then the particulars of the columnar pieces of FS might be important. The changing sizes of these pieces with $x$ in the ARPES data (but not the dHvA data) would be consistent with this idea.

In conclusion, while good progress has been made in determining the electronic structure of this interesting new alloy series, it remains to find a unified view that explains both what is now known about the electronic properties and what is known about the transport properties. One step forward for the future would be to obtain a complete set of ARPES data for intermediate values of $x$, ideally having the same high quality as that reported here for $x = 0$ and $x = 1$.

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