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Zn-flux-enabled synthesis of orthorhombic kagome YbFe_6Ge_6 : Yb reduction and magnetic behavior

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Single crystals of an orthorhombic polymorph of kagome metal YbFe_6Ge_6 were grown from Zn flux. This Zn-flux-stabilized phase exhibits mixed Yb valency and magnetic behavior that differs from its hexagonal counterpart, underscoring the profound influence of flux selection on both structure and physical properties in intermetallic materials.

Flux growth synthesis is an efficient method for producing bulk single-crystalline materials.^{1,2} The technique not only enables the discovery of compounds within complex phase spaces, but also facilitates the growth of large single crystals suitable for physical measurements. Flux growth leverages eutectic points in a mixture of elements, often utilizing a low-melting metal or compound to reduce the overall melting point of a reaction. The technique permits synthesis with refractory elements to be more accessible, thus making it particularly valuable for the exploratory growth of intermetallic phases. Conventional synthesis methods for intermetallic compounds typically promote the formation of thermodynamically stable phases within a given composition space, thereby limiting access to metastable or otherwise unexplored structural regions that require kinetic or non-equilibrium control.³ Flux growth can circumvent this by altering reaction pathways through adjustable experimental parameters intrinsic to the technique. For example, the formation of specific products can be guided by tuning the cooling rate or by selecting the temperature at which the reaction is removed from the furnace and the excess flux is removed. This level of control was demonstrated through the investigation of the Ce–Pd–Ga phase space where the use excess Ga flux led to the discovery of a family of structurally related compounds.^{2,4} A

range of related structures were obtained by tuning the heating parameters, which illustrate the diverse parametric control of flux growth methods. In the synthesis of the $\text{Pr}_{n+1}\text{Co}_n\text{Ge}_{3n+1}$ homologous series,⁵ the concentration of flux used directly affected which n -members and related phases crystallized from the melt. This study reinforced the notion that flux concentration itself serves as a valuable tuning parameter for directing phase formation.⁶ Harnessing control over structural formation within a phase space is also evident in the stabilization of the two polymorphs of $\text{Pr}_2\text{Co}_3\text{Ge}_5$, one orthorhombic⁷ and one monoclinic, the latter displaying nearly Pr^{4+} character, which is rare for Pr intermetallic compounds.⁸

In this communication, we demonstrate how the choice of flux can decisively influence phase formation in intermetallic synthesis. YbFe_6Ge_6 is a member of the broader family of rare-earth kagome metals.^{9,10} The HfFe_6Ge_6 -type structure ($h\text{-YbFe}_6\text{Ge}_6$, SG: $P6/mmm$) is routinely grown from a Sn flux and hosts a low temperature spin reorientation near 63 K,^{11,12} a feature associated with an anomalous Hall effect.¹³ The $\text{Y}_{0.5}\text{Co}_3\text{Ge}_3$ -type structure ($y\text{-YbFe}_6\text{Ge}_6$, SG: $P6/mmm$) can also be obtained through conventional arc-melting and annealing at 900 °C and does not possess the spin reorientation present in the ordered variant.¹⁴ Here, we show that loading Yb, Fe, Ge with an excess of Zn flux, yields an orthorhombic polymorph of YbFe_6Ge_6 ($o\text{-YbFe}_6\text{Ge}_6$, SG: $Cmcm$) that adopts the TbFe_6Sn_6 structure type and, exhibits differing magnetic properties than its hexagonal counterpart. Energy-dispersive X-ray spectroscopy confirms Zn incorporation into the structure while powder diffraction indicates a preferential Zn site occupation within the Fe-based kagome slabs (Fig. S2). For simplicity, we refer to Zn-flux-stabilized orthorhombic phase, $\text{YbFe}_{5.52}\text{Zn}_{0.48}\text{Ge}_6$, as $o\text{-YbFe}_6\text{Ge}_6$ throughout this communication. Single crystals of $o\text{-YbFe}_6\text{Ge}_6$ were grown out of excess Zn flux with sizes up to 5 mm, adopting a hexagonal prism morphology (Fig. S1, inset). Synthesis conditions and structure determination using powder X-ray diffraction, single crystal X-ray diffraction and energy dispersive X-ray spectroscopy are provided in the SI.

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Fig. 1 shows the unit cell of o -YbFe₆Ge₆ (a) and the patterning scheme of the Yb sublattice (b). Crystallographic data and refinement parameters for o -YbFe₆Ge₆ are given in Table S2 and atomic coordinates of o -YbFe₆Ge₆ are given in Table S3. Crystallographic comparison with h -YbFe₆Ge₆ was performed using the CIF from our previous report, as the reactant stoichiometry offers a more meaningful comparison to the orthorhombic variant under similar synthesis conditions.¹⁵ Both the h -YbFe₆Ge₆ and o -YbFe₆Ge₆ structures consist of alternating honeycomb and kagome slabs that are stacked in a staggered fashion, forming channels which house the Yb and the Ge–Ge dimer (Fig. S3a and b). In h -YbFe₆Ge₆, Yb occupies every other honeycomb slab along the c -axis. Structural complexity increases in o -YbFe₆Ge₆ due to differences in Yb ordering, which represents the main feature of distinction between the two structure types. Assuming the absence of the disordered sites, in h -YbFe₆Ge₆, the Yb atoms centered within the honeycomb hexagons are adjacent to one another in the ab plane. In contrast, o -YbFe₆Ge₆ exhibits a staggered Yb arrangement along the b -axis, such that every other pair of Yb atoms is adjacent. Unlike h -YbFe₆Ge₆, where Yb occupies every other honeycomb layer along the c -axis, the orthorhombic variant possesses Yb in each honeycomb layer along the a -axis, resulting in a more intricate ordering scheme. The reason behind the specific ordering was discussed in detail by Fredrickson *et al.*¹⁶ It is seen that with larger rare earth elements occupying the hexagons of the honeycomb layer (or other constituent elements that surround the rare earth), they are increasingly unable to order adjacent to one another because of increased steric interactions, leading to a staggered ordering of the rare

earth element. One detour from the hexagonal to orthorhombic transition is seen with the hexagonal Yb_{0.5}Co₃Ge₃ structure type; that harbors large amounts of positional disorder of Yb and Ge dimers, resulting in a reduction of the hexagonal HfFe₆Ge₆-type cell by $c/2$. Our previous report of Yb_{0.5}(Co_{1-x}Fe_x)₃Ge₃ explores the structural transition from a Yb_{0.5}Co₃Ge₃/CoSn hybrid structure to the HfFe₆Ge₆ structure type as a function of Fe substitution.¹⁵

Fig. S4c and d show the honeycomb sublattice of the h -YbFe₆Ge₆ and o -YbFe₆Ge₆ structures, respectively with distinct interatomic distances highlighted to show how the perfect hexagonal slab in the h -YbFe₆Ge₆ structure differs from the irregular hexagons present in o -YbFe₆Ge₆. Fig. S4e and Fig. S4f show the areas of the h -YbFe₆Ge₆ and o -YbFe₆Ge₆ hexagons, respectively. Fig. S3c and d show the patterning scheme of the Ln environment in h -YbFe₆Ge₆ and o -YbFe₆Ge₆ which differs based on the occupancy of the lanthanide in the honeycomb net. Using the rationale that unfavorable steric environments can drive an orthorhombic distortion, we propose that Zn incorporation on the Fe sites reduces the local volume surrounding the Yb atoms in a manner reminiscent trends in of our previous doping study.¹⁵ Another possibility we considered is that Yb in o -YbFe₆Ge₆ may not be strictly trivalent; partial Yb²⁺ character could increase the average ionic radius and therefore influence the observed structural distortion. We also note that differences in synthesis conditions between h -YbFe₆Ge₆ and o -YbFe₆Ge₆ may play a role. Synthesis-dependent polymorphism has been documented in YFe₆Sn₆ and DyFe₆Sn₆,¹⁷ and studies of FeGe also previously showed that the annealing conditions affected the structural ordering,¹⁸ thus in our Yb_{0.5}(Co_{1-x}Fe_x)₃Ge₃ report, an annealing step of 72 h was performed to ensure structural consistency in the doped series; such post-growth annealing was not performed here. When attempting to use the same heating profile as our previous report, the orthorhombic polymorph consistently forms with the only difference in structure being the amount of positional disorder present, which could be due to either the change in cooling rate or the time dwelled.

X-ray absorption spectroscopy (XAS) performed at the Yb L_{3-} edge was used to quantify the Yb oxidation state in o -YbFe₆Ge₆ (see SI for measurement details). The resulting normalized spectra for o -YbFe₆Ge₆ and the known +3 oxidation state reference YbCl₃ are shown in Fig. 2. Measurement of the edge energy *via* derivative analysis (Fig. S5c), reveals a small redshift in the edge energy from 8944.11 eV for YbCl₃ to 8942.68 eV for o -YbFe₆Ge₆ ($\Delta E = -1.43$ eV) consistent with an oxidation state assignment Yb³⁺ for the YbFe₆Ge₆ analyte.¹⁹ In addition to this main feature, the derivative of the spectrum displayed an additional, lower energy feature that contributes to the broad nature of the primary peak. Peak fitting analysis (see SI) was performed to deconvolute the spectrum. This revealed a secondary, lower intensity peak centered at 8938.27(5) eV, redshifted ~ 6.6 eV from the main peak at 8944.846(16) eV. While the higher energy peak is consistent with Yb³⁺, the lower energy peak is consistent with a small percentage of Yb²⁺ character in the compound and this intermediate-valence effect has been

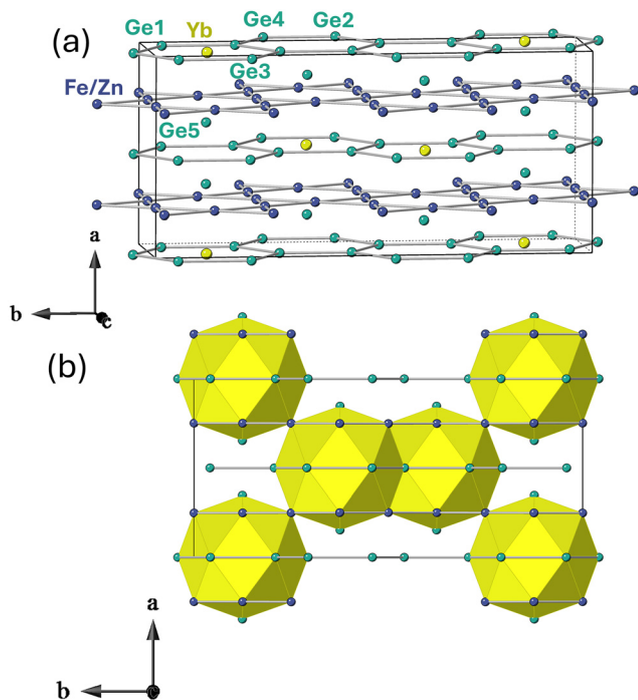


Fig. 1 Unit cell (a) and patterning scheme (b) along the b -direction of o -YbFe₆Ge₆, disordered sites omitted for clarity.



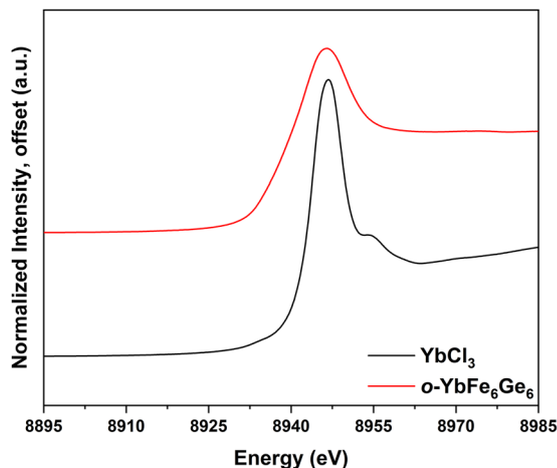


Fig. 2 Normalized (offset) Yb L_3 -edge spectra for YbCl_3 (black) and $o\text{-YbFe}_6\text{Ge}_6$ (red).

observed in other Yb-containing intermetallic compounds.^{20–23} The appearance of Yb^{2+} character in these systems is associated with coupling of the conduction band electrons to the $4f$ hole of Yb^{3+} , which itself lies near the Fermi energy. The increase of valency in the kagome sublattice with Zn substitution can also be contributing to the reduction of Yb. To estimate the relative $\text{Yb}^{2+/3+}$ character of $o\text{-YbFe}_6\text{Ge}_6$, the ratio of the peak area of the Yb^{3+} feature to the sum of the area of both peaks, denoted n_f (eqn (S1)), is calculated and found to be 0.79(8), which is consistent with the assignment of a formal Yb^{3+} oxidation state, with relatively minor amounts of Yb^{2+} character and an effective oxidation state (eqn (S2)) of +2.79(8) at the measurement temperature of 7 K.

Fig. 3a and b presents the anisotropic temperature-dependent magnetic susceptibility measurements when the applied

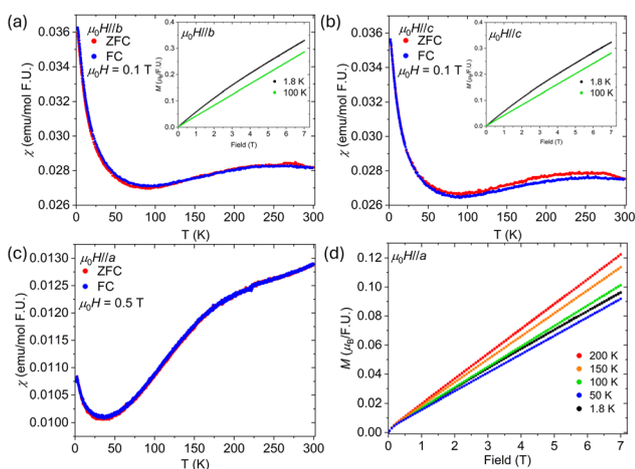


Fig. 3 Temperature-dependent magnetic susceptibility of $o\text{-YbFe}_6\text{Ge}_6$ where (a) $\mu_0H\parallel b$ and (b) $\mu_0H\parallel c$. The inset of both plots is the field-dependent magnetization data up to 7 T at 100 K and 1.8 K isotherms. (c) Temperature-dependent magnetic susceptibility where $\mu_0H\parallel a$. (d) Field-dependent magnetization up to 7 T at 200 K, 150 K, 100 K, 50 K, and 1.8 K where $\mu_0H\parallel a$.

magnetic field is parallel to the b -direction and c -direction ($\mu_0H\parallel b$ and $\mu_0H\parallel c$), respectively. The data for both field orientations are largely comparable, showing a broad hump beginning near 300 K, which suggests possible spin canting or fluctuations of the Fe moments down to approximately 100 K, where the emergence of Yb paramagnetism becomes apparent. The magnetization *versus* applied magnetic field measurements (inset) at $T = 100$ K and $T = 1.8$ K also exhibit similar behavior, characterized by an almost linear field dependence. These similarities for fields applied parallel to the b - and c -directions can be attributed to the fact that in both cases the field lies parallel to the kagome planes. Fig. 3c and d present the temperature-dependent magnetic susceptibility and the field-dependent magnetization, respectively, when the field is applied parallel to the a -direction ($\mu_0H\parallel a$). The susceptibility exhibits a gradual decrease from 300 K to about 50 K where a paramagnetic upturn is observed. Field-dependent magnetization measurements were collected at several temperatures to examine how changes in magnetic field strength influence the downturn observed in the magnetic susceptibility. The 200 K isotherm exhibits the largest magnetization across the measured field range. As the temperature decreases, the overall magnetization systematically diminishes until 50 K, where the onset of the Yb paramagnetic contribution becomes apparent, consistent with the susceptibility data. At 1.8 K, the magnetization increases slightly relative to the 50 K isotherm, reflecting the growing Yb paramagnetism at low temperatures. LnFe_6Ge_6 members ($\text{Ln} = \text{Gd-Lu}$) adopt an antiferromagnetic ordering of Fe moments along the kagome stacking axis at approximately 480–490 K.⁹ In compounds adopting the HfFe_6Ge_6 structure type, this axis corresponds to the c -axis, whereas in those with the TbFe_6Sn_6 structure type, it corresponds to the a -axis (Fig. 1). Therefore, in $o\text{-YbFe}_6\text{Ge}_6$ (TbFe_6Sn_6 -type), the a -axis can be considered the magnetic “hard axis,” consistent with the orientation of the antiferromagnetically ordered Fe moments. Direct confirmation of the Fe-sublattice magnetic transition would require high-temperature magnetic susceptibility measurements, as the transition is expected to occur well above room temperature. This interpretation is further supported by the reduced magnitude of the susceptibility for $\mu_0H\parallel a$ compared to $\mu_0H\parallel b$ and $\mu_0H\parallel c$. Reliable Curie–Weiss fits could not be obtained because the Fe sublattice likely orders above 300 K, preventing access to a true paramagnetic regime for the Fe moments. The paramagnetic upturn observed below ~ 50 K is attributed to the emergence of localized Yb^{3+} moments, which begin to dominate the magnetic response as thermal fluctuations decrease. At higher temperatures, the broad behavior of the magnetic susceptibility is governed primarily by Fe-based interactions, potentially involving weak spin canting or short-range correlations within the kagome planes. This is most likely a result of Zn incorporation into the Fe lattice, which diminishes the long-range ordering normally exhibited in a pure Fe kagome slab. As the temperature decreases, the Yb^{3+} ions display Curie–Weiss-like behavior, producing the rapid increase in susceptibility. This temperature evolution reflects a crossover from itinerant Fe magnetism at high temperatures to localized rare-earth paramagnetism at low temperatures,



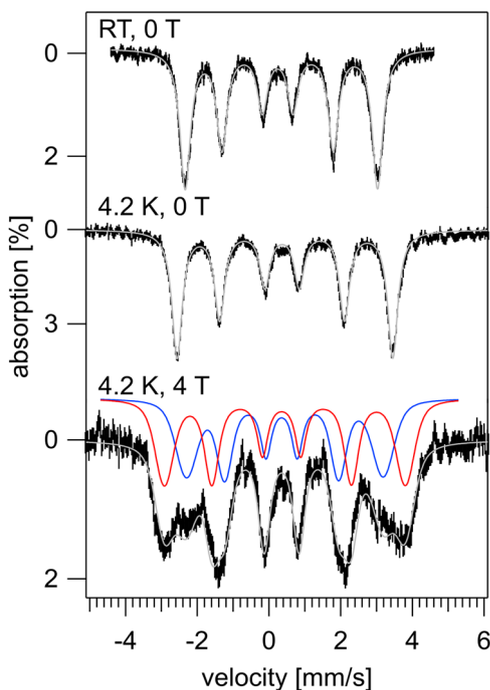


Fig. 4 ^{57}Fe Mössbauer spectra recorded for $o\text{-YbFe}_6\text{Ge}_6$, see text. The solid grey traces are theoretical spectra obtained using the parameters listed in Table S4. The theoretical 4 T spectrum was obtained from the sum of two components shown in blue and red above the experimental data.

consistent with the onset of Yb-moment polarization in applied magnetic fields.

The temperature dependent, zero-field ^{57}Fe Mössbauer spectra recorded for $o\text{-YbFe}_6\text{Ge}_6$ exhibit a single well defined six-line pattern, see Fig. 4 and Fig. S7. These spectra demonstrate that there is a spontaneous magnetic field acting on the ^{57}Fe nuclei which reveals that the iron sublattice is magnetically ordered. Inspection of Fig. 4, Fig. S8, and Table S4 shows that the hyperfine field increases from $B_{\text{eff}} = 16.6$ T at room temperature to 18.6 T at 4.2 K. These values are $\sim 10\%$ larger than those reported for $h\text{-}$ and $y\text{-}$ type YbFe_6Ge_6 .^{11,14,24,25} Assuming that the 11.39 T μ_{B}^{-1} conversion factor determined at 150 K for $h\text{-YbFe}_6\text{Ge}_6$ from its combined neutron diffraction and ^{57}Fe Mössbauer spectroscopic investigation is still valid, these values suggests that for $o\text{-YbFe}_6\text{Ge}_6$ the iron-based magnetic moment increases from $1.46\mu_{\text{B}}$ at room temperature to $1.63\mu_{\text{B}}$ at 4.2 K. Importantly, lowering the temperature below 80 K does not lead to a broadening or a splitting of the observed resonances. This indicates that, just like for $y\text{-YbFe}_6\text{Ge}_6$, the spin reorientation transition observed at ~ 63 K for $h\text{-YbFe}_6\text{Ge}_6$ is suppressed by the increased disorder of the Yb sites. Field dependent data up to 8 T was also collected (Fig. S9). These spectra suggest that above 6 T, the onset of a metamagnetic transition may occur even with no obvious features in the magnetization.

Several factors may account for the differences in magnetic behavior. (1) In the $h\text{-YbFe}_6\text{Ge}_6$ polymorph, XAS measurements confirm that Yb is trivalent. However, it is seen here that the $o\text{-YbFe}_6\text{Ge}_6$ polymorph has mixed valency that could potentially

alter Yb–Fe coupling. (2) Partial substitution of Fe by Zn within the kagome lattice suppresses magnetic exchange among Fe atoms, thereby weakening the Yb–Fe coupling through alloy dilution. (3) The magnetic distinctions may also arise from differences in the structural ordering of the Yb sublattice. $h\text{-YbFe}_6\text{Ge}_6$ features a well-ordered triangular arrangement of Yb atoms, while the highly disordered $y\text{-YbFe}_6\text{Ge}_6$ contains a random Yb distribution and is reported to lack the spin reorientation present in the ordered variant.¹⁴ The specific patterning scheme of Yb atoms we observe in $o\text{-YbFe}_6\text{Ge}_6$ may similarly disrupt Yb-Fe exchange interactions. Given that the temperature dependent Mössbauer spectra manifests similar behavior as the $y\text{-YbFe}_6\text{Ge}_6$, the origin in the differences in magnetism seem to strongly point to the pattern of the Yb sublattice. In conclusion, the reduction of Yb from a Zn flux growth leads to a change in physical properties and structure in $o\text{-YbFe}_6\text{Ge}_6$ when compared to the previously reported $h\text{-YbFe}_6\text{Ge}_6$. Clarifying the mechanism of growth and its influence on the contrasting physical properties of $h\text{-YbFe}_6\text{Ge}_6$ and $o\text{-YbFe}_6\text{Ge}_6$ will require the use of *in situ* diffraction and spectroscopic methods and are a prospect of future work. The present study underscores the critical role of parametric control of the flux growth method as well as its significance in enabling the synthesis of previously inaccessible phases.

Conflicts of interest

There are no conflicts to declare.

Data availability

The data supporting this article are included as part of the supplementary information (SI). Supplementary information: additional experimental details, XRD, crystallographic data, XAS analysis, and additional Mössbauer spectra. See DOI: <https://doi.org/10.1039/d6cc00958a>.

CCDC 2528289 contains the supplementary crystallographic data for this paper.²⁶

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