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Ternary borides Nb₇Fe₃B₈ and Ta₇Fe₃B₈ with Kagome-type iron framework†

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Two new ternary borides $TM_7Fe_3B_8$ (TM = Nb, Ta) were synthesized by high-temperature thermal treatment of samples obtained by arc-melting. This new type of structure with space group P6/mmm, comprises TM slabs containing isolated planar hexagonal $[B_6]$ rings and iron centered TM columns in a Kagome type of arrangement. Chemical bonding analysis in $Nb_7Fe_3B_8$ by means of the electron localizability approach reveals two-center interactions forming the Kagome net of Fe and embedded B, while weaker multicenter bonding present between this net and Nb atoms. Magnetic susceptibility measurements reveal antiferromagnetic order below $T_N = 240$ K for $Nb_7Fe_3B_8$ and $T_N = 265$ K for $Ta_7Fe_3B_8$. Small remnant magnetization below $0.01\mu_B$ per f.u. is observed in the antiferromagnetic state. The bulk nature of the magnetic transistions was confirmed by the hyperfine splitting of the Mössbauer spectra, the sizable anomalies in the specific heat capacity, and the kinks in the resistivity curves. The high-field paramagnetic susceptibilities fitted by the Curie–Weiss law show effective paramagnetic moments $\mu_{eff} \approx 3.1\mu_B/Fe$ in both compounds. The temperature dependence of the electrical resistivity also reveals metallic character of both compounds. Density functional calculations corroborate the metallic behaviour of both compounds and demonstrate the formation of a sizable local magnetic moment on the Fe-sites. They indicate the presence of both antiferro- and ferrromagnetic interactions.

Introduction

Transition-metal borides are remarkable for their physical, chemical and mechanical properties, in particular, combining refractory behavior, high hardness, chemical inertness, and metallic conductivity. To exemplify a few well-known materials of high technological relevance, there are boride-containing metallic glasses, LaB₆ cathodes for electron microscopes, and Nd₂Fe₁₄B-based permanent magnets. On the other hand, soft magnetic properties particularly found in ferrous amorphous boron-containing alloys have led to several

highly useful applications, such as electro-magnetic materials. The structural complexity of electron-deficient boron and its compounds is caused by the intricate ways how their valence requirements are satisfied. Therefore, in metal-borides this not only gives rise to the formation of one-, two-, or three-dimensional arrangements of covalently bonded boron atoms, the due to complex structures, results in a multitude of physical interactions which can lead to superconductivity 22-26 or magnetism. 27-31

In a crystal structure, these interactions are governed by the spatial arrangement and coordination of the constituents which carry a magnetic moment. Accordingly, in borides phases with exotic and complex magnetic ground states can be expected and merit explorative research. To this purpose, we investigated the ternary TM–Fe–B (TM = Nb, Ta) systems which have been studied since the 1960s. The hitherto known ternary compounds in these two systems are TMFeB, TM_2FeB_2 , $TM_3Fe_3B_4$ and $TaFeB_3$. We attempted to synthesize $TM_3Fe_3B_4$ but failed. Instead, the analysis of samples with nominal compositions $TM_3Fe_3B_4$ revealed the appearance of new compounds $TM_7Fe_3B_8$, which crystallize in a primitive hexagonal lattice. This new type of structure comprises TM slabs containing planar hexagonal boron rings and iron centered TM columns in a Kagome type of arrangement (see below).

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The interest in the Kagome lattices of magnetic ions is triggered by their strongly frustrated nature. No ordered antiferromagnetic configuration can be stabilized in such a geometry, and an exotic spin-liquid ground state is formed instead. 36,37 However, in order to suppress any ordering a two-dimensional isotropic next-neighbor coupling is the prerequisite. Systems with the Kagome-like arrangement of magnetic ions range from Cu²⁺ minerals^{38,39} to Ce-based intermetallic compounds, 40 but only few of them reveal the anticipated spinliquid state at low temperatures. 41-43 Even subtle geometrical distortions or interactions beyond nearest neighbors are usually sufficient to alleviate the frustration and stabilize the magnetic order.44,45

The majority of the kagome-lattice compounds reported so far are magnetic insulators. Kagome lattices in itinerant systems are by far more exotic, and no magnetic metals with the ideal kagome geometry have been reported to date. In the following, we fill this gap by investigating the TM7Fe3B8 borides, where three-fold symmetry of the crystal structure ensures perfect frustration on individual triangles of the kagome lattice. However, these compounds are magnetically ordered with relatively high Néel temperatures. We suggest that strong interplane couplings arising from the geometrical proximity of the kagome planes may be instrumental in stabilizing the magnetically ordered state in these novel ternary Fecontaining borides.

Experimental section

Sample preparation

The following elements were used to prepare the samples: Ta and Nb foil (Chempur, 99.9 mass%), Fe foil (Alfa Aesar, 99.999 mass%) and crystalline B powder (Alfa Aesar, 99.999 mass%). The mixtures of the elements with compositions TM₃Fe₃B₄ and TM₇Fe₃B₈ were arc-melted several times to obtain homogeneous samples, and the mass losses during arc-melting were 1-3.3%. The obtained ingots were placed inside Al₂O₃ crucibles and then sealed in Ta tubes. The annealing was carried out at 1000 °C for 12 h and then 1500 °C for 48 h in a high-temperature furnace (HTM Reetz, LORA). Additionally, all above manipulations were performed inside argon-filled glove boxes $(p(O_2/H_2O) \le 1 \text{ ppm})$. The resulting samples are stable in air for a long time.

Powder and single-crystal X-ray diffraction

Powder X-ray diffraction (XRD) data were collected on a HUBER G670 imaging plate Guinier camera equipped with Co $K_{\alpha 1}$ radiation ($\lambda = 1.78897$ Å). Phase analysis and indexing were performed using the WinXPow program package. 46 Lattice parameters were refined by least-squares fitting with LaB₆ internal standard within the program package WinCSD.⁴⁷

TM₇Fe₃B₈ single crystals were selected from the samples with the nominal compositions TM₃Fe₃B₄. Single crystal diffraction data were collected on a R-Axis Rapid or Rigaku AFC7 diffractometer equipped with Mercury CCD detectors (Mo K_o

Table 1 Crystallographic data for Nb₇Fe₇B₈ and Ta₇Fe₇B₈

Composition	$\mathrm{Nb_{7}Fe_{3}B_{8}}$	$\mathrm{Ta_{7}Fe_{3}B_{8}}$
Crystal system	hexagonal	
Space group	<i>P</i> 6/ <i>mmm</i> (no. 191)	
Lattice parameters	` ,	
a (Å)	8.3346(2)	8.2788(3)
c (Å)	3.2941(1)	3.2934(3)
$V(A^3)$	198.17(2)	195.48(3)
Formula unit/cell, Z	1	1
Calculated density	7.5777(6)	12.916(2)
(g cm ⁻³)		
Crystal size (mm ³)	$0.02\times0.02\times0.02$	$0.04\times0.04\times0.04$
Diffraction system	Rigaku R-Axis Rapid	Rigaku AFC7
Radiation; λ (Å)	Mo $K_α$; 0.71073 Å	
$2\theta_{\mathrm{max}}$ (°)	131.69	66.40
Absorption coefficient	14.751	108.146
(mm^{-1})		
$N(hkl)_{\text{measured}}$	5932	1774
N(hkl) _{unique}	719	179
$N(hkl)_{\text{observed}} (F_{hkl} > 4\sigma(F))$	614	174
$R_{ m int}/R_{\sigma}$	0.050/0.023	0.045/0.016
Refined parameters	17	14
R_F/WR_F^2	0.047/0.053	0.017/0.017
Extinction coefficient	0.063(4)	0.026(1)

radiation, $\lambda = 0.71073$ Å). Absorption correction was made using a multi-scan procedure. The crystal structures were solved by a direct phase determination method and refined by a full-matrix least-squares procedure within the program package WinCSD.47 Details on the single-crystal diffraction data collection and structural refinement are listed in Table 1.

Metallography

Pieces with several millimeters size were cut from the annealed samples for metallographic investigations. They were embedded in conductive resin and then subjected to a multistep grinding and polishing process to achieve high-quality polished surfaces. The microstructures were investigated by optical microscopy (Axioplan2, Zeiss) as well as scanning electron microscopy (Philips XL 30 with a LaB₆ cathode, FEI). The chemical compositions were analyzed by means of energy dispersive X-ray spectroscopy (EDXS, Philips XL 30) and wavelength dispersive X-ray spectroscopy (WDXS, Cameca SX 100, W cathode, S-UTW-Si-(Li) detector).

The determination of the boron content by microprobe WDXS is challenging due to the general issue that the measured intensities are related to the mass concentrations, while boron is extremely light as compared to niobium, tantalum, and iron. Also, the very low energies of boron X-ray lines give rise to a strong influence of absorption effects. Therefore, completely detected intensities originating from the uppermost surface layer and its extending area are strongly influenced by the quality of the local area that is probed by the electron beam, and the energy and shape of the boron X-ray lines are influenced significantly by the local chemical environment and bonding situation of boron. For this reason, reference compounds should be materials, in which boron should have a similar chemical environment and bonding situation with the two analyzed compounds. Subsequently, single

crystals of NbFeB, TaFeB in sizes of several millimeters were successfully grown in melt fluxes and satisfy requirement for WDXS reference materials. By applying ICP-MS technique, their analyzed compositions are Nb_{1.00(1)}Fe_{1.02(1)}B_{1.01(2)} and Ta_{1.00(1)}Fe_{1.03(1)}B_{1.01(1)}, respectively, revealing both are stoichiometric.⁴⁸ The intensities of Ta L_{α} , Fe K_{α} and B K_{α} lines were measured for the Ta-containing compound, while due to the overlapping of B K_{α} line with Nb M_{ν} line, the intensities of Nb L_{α} , Fe K_{α} and B $K_{\alpha 2}$ lines were measured for the Nb-containing compound.

The matrix correction model according to Pouchou and Pichoir⁴⁹ was applied to calculate the chemical compositions. Different conditions have been applied for the measurement of the X-ray lines of the heavy elements and boron. For the Tacontaining compound, currents of 15 nA and 40 nA under an acceleration voltage of 20 kV were applied for the measurement of the intensities of Ta L_{α} and Fe K_{α} lines, respectively, with the dwelling time of 3 seconds for each position. The intensity of the boron K_{α} line was measured by the area intensity method, with larger current of 100 nA, acceleration voltage of 7 kV and dwelling time 3 seconds for each position. For the Nb compound, the intensities of Nb L_{α} and Fe K_{α} lines were measured by applying a current of 60 nA under an acceleration voltage of 15 kV with dwelling time 3 seconds for each position. The intensity of the B $K_{\alpha 2}$ line was measured by applying the same acceleration voltage and current used for the Ta compound, however, due to the much weaker intensity of the B $K_{\alpha 2}$ line, dwelling time for each position was 1136 seconds.

Transmission electron microscopy (TEM) observations

Electron diffraction and high-resolution TEM (HRTEM) observations were both performed using a field-emission electron microscope JEM 2100F (JEOL, Japan) operating at 200 kV. HRTEM image simulations were carried out with program STEM CELL.50

Physical properties

Magnetization at external magnetic fields $\mu_0 H$ ranging from 0.01 T to 7 T (temperature range 1.8 K-400 K) was measured in a SQUID magnetometer (MPMS XL-7, Quantum Design) on polycrystalline samples. The electrical resistance was recorded by a four contact method using low-frequency alternating current (ACT option, PPMS, Quantum Design) on small barshapes pieces in zero field and in a field $\mu_0 H = 9$ T. Heat capacity was determined by a relaxation method (HC option, PPMS, Quantum Design) in fields $\mu_0 H$ of 0, 3, 6, and 9 T.

Mössbauer spectroscopy

⁵⁷Fe Mössbauer spectra were recorded at 294 K and 4.3 K. The measurements were performed with a standard constant acceleration spectrometer in transmission geometry in a continuous flow cryostat with the sample kept at helium atmosphere. The ⁵⁷CoRh source was mounted on the driving system and kept at room temperature. All center shift (CS) data are given relative to this source. Calibration of the velocity scale was carried out with α-Fe foils. The spectra were analysed by solving the full Hamiltonian including electrostatic and magnetic hyperfine interactions. Sample thickness was taken into account by the method of Mørup and Both.⁵¹

Electronic structure calculations

The electronic structure of TM₇Fe₃B₈ was calculated within the framework of density functional theory (DFT) using the fullpotential code FPLO.⁵² The local density approximation (LDA) to the exchange-correlation potential was chosen. 53 Reciprocal space was sampled by a fine k-mesh with 630 points in the symmetry-irreducible part of the first Brillouin zone for the crystallographic unit cell of TM7Fe3B8 and 190 points for the supercell doubled along the c direction. Convergence with respect to the k-mesh was carefully checked. For the spinpolarized calculations, the highest crystallographic symmetry compatible with the magnetic ordering pattern was used in order to facilitate the convergence.

Chemical bonding analysis

Analysis of chemical bonding was performed for Nb₇Fe₃B₈ using the lattice parameters and atomic coordinates from the crystal structure refinement of single-crystal X-ray diffraction data (Tables 1 and 2). The TB-LMTO-ASA program package⁵⁴ was employed using the Barth-Hedin exchange potential⁵⁵ for the LDA calculations. The radial scalar-relativistic Dirac equation was solved to obtain the partial waves. 56 Addition of empty spheres was not necessary because the calculation within the atomic sphere approximation (ASA) includes corrections for the neglect of interstitial regions and partial waves of higher order.⁵⁷ The following radii of the atomic spheres were applied for the calculations r(Nb1) = 1.69 Å, r(Nb2) = 1.49 Å,r(Fe) = 1.48 Å, r(B1) = 1.02 Å, r(B2) = 1.19 Å. A basis set containing Nb(5s,5p,4d), Fe(4s,4p,3d) and B(2s,2p) orbitals was employed with Nb(4f) and B(3d) functions being downfolded.

The electron localizability indicator (ELI, Y) was evaluated in the ELI-D representation⁵⁸⁻⁶⁰ with an ELI-D module within the TB-LMTO-ASA program package. Topological analysis of the electron localizability indicator, e.g., localization of the ELI maxima as indicators of the direct atomic interactions, estimation of their basins were performed with the program DGrid.61

Table 2 Atomic coordinates and isotropic displacement parameters for Nb₇Fe₃B₈ and Ta₇Fe₃B₈

Atom	Site	x	у	z	$B_{\rm iso}/B_{\rm eq}^{a}$
Nb ₇ Fe ₃ B ₈					
Nb1	1a	0	0	0	0.30(2)
Nb2	6l	0.21202(3)	2x	0	0.313(8)
Fe	3g	1/2	0	1/2	0.41(2)
B1	2d	1/3	2/3	1/2	0.5(1)
B2	6k	0.2186(8)	0	1/2	0.51(9)
Ta ₇ Fe ₃ B ₈					
Ta1	1a	0	0	0	0.20(2)
Ta2	6l	0.21233(3)	2x	0	0.27(2)
Fe	3g	1/2	0	1/2	0.34(5)
B1	2d	1/3	2/3	1/2	0.3(1)
B2	6k	0.221(1)	0	1/2	0.4(1)

 $^{^{}a}$ $B_{\text{eq}} = \frac{1}{3} \left[a^{*2} a^{2} B_{11} + b^{*2} b^{2} B_{22} + c^{*2} c^{2} B_{33} + 2aba^{*} b^{*} (\cos \gamma) B_{12} + 2aca^{*} c^{*} \right]$ $(\cos \beta)B_{13} + 2bcb*c*(\cos \alpha)B_{23}$].

Results and discussion

Phase formation and crystal structure determination

Both as-cast samples with the nominal compositions TM₃Fe₃B₄ and TM₇Fe₃B₈ contain Fe₂B, TMB and a little amount of other TM-B binary phases, however, with no traces of any other ternary phases. The TM7Fe3B8 compounds were firstly discovered as matrix phases in the samples with nominal composition TM₃Fe₃B₄ annealed at 1500 °C for 48 h. Subsequently, bulk samples of TM₇Fe₃B₈ with minor amounts of impurity phases were obtained by annealing at 1500 °C for 48 h. However, both compounds decomposed into TMB, Fe₂B phases (and sometimes with α -Fe) at temperatures of about 1600 °C. The decomposition was proved by an endothermal peak in DSC curves at ≈ 1580 °C for the Nb₇Fe₃B₈ sample.

Powder XRD patterns of the sample TM7Fe3B8 annealed at 1500 °C are shown in Fig. 1. 26 strongest reflections in the Nbcontaining sample were indexed successfully in a hexagonal primitive lattice with unit cell parameters a = 8.3346(2) Å, c =3.2941(1) Å (Figure of merit (FOM) = 72.7) using the automatic indexing algorithm TREOR within WinXPow program package. In the same way, 23 strongest reflections in the Ta-containing sample were also successfully indexed in a hexagonal primitive unit cell with lattice parameters a = 8.2788(3) Å, c = 3.2934(3) Å (FOM = 78.2). No extinction conditions were observed in the two data sets. The remaining reflections belong to the TMB phases (\sim 1 vol% and \sim 3 vol% in Nb₇Fe₃B₈ and Ta₇Fe₃B₈, respectively).

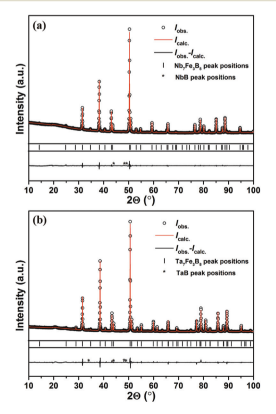


Fig. 1 Powder XRD patterns of the samples with nominal composition (a) Nb₇Fe₃B₈ and (b) Ta₇Fe₃B₈ annealed at 1500 °C.

The results of the single-crystal X-ray diffraction experiment are well consistent with the powder pattern indexing. Analysis of the reflection conditions revealed Laue class 6/mmm, and the most symmetric space group P6/mmm was chosen to solve the crystal structures. Since light B atoms give only a small contribution to the reflection intensities, reflexes were measured up to high angles ($2\theta_{\text{max}} = 131.69^{\circ}$) for the Nb-containing single crystal in order to facilitate the determination of the positions of B atoms. The initial positions of Nb/Ta and Fe atoms were obtained by the direct phase determination method, while the positions of B atoms were found from difference Fourier maps. The refinement was carried out with anisotropic approximation for the atomic displacement parameters for all the atoms (except B atoms in Ta₇Fe₃B₈ where refinement due to relatively large absorption is questionable). Occupations of the Ta, Nb and Fe positions were also refined, revealing that all are fully occupied. Crystallographic data and final atomic coordinates as well as isotropic displacement parameters are listed in Tables 1 and 2. Anisotropic displacement parameters are deposited in Table S1 in the ESI.† Compositions are in good agreement with WDXS results. By WDXS, the compositions for the two compounds were achieved as follows: $Ta_{7.00(5)}Fe_{3.13(3)}B_{7.92(7)}$ and $Nb_{7.00(6)}Fe_{3.10(3)}B_{7.4(2)}$. They are satisfactorily close to the compositions TM₇Fe₃B₈ obtained from single crystal diffraction data, and the former one is nearly the same. The lower precision and accuracy of the obtained boron content for the Nb-containing compound results mainly from using the much weaker B $K_{\alpha 2}$ line. Moreover, WDXS compositions are consistent with the results by standardless EDXS analysis, which yield the Ta: Fe and Nb: Fe atomic ratios 70.6(8): 29.4(8) and 70(1): 30(1), respectively.

TEM investigation

The TEM study of Ta₇Fe₃B₈ confirmed the results of the crystal structure determined by the X-ray diffraction. The electron diffraction patterns along relevant zone axes are shown in Fig. 2. All five patterns can be well indexed in a hexagonal primitive lattice with the cell parameters obtained from

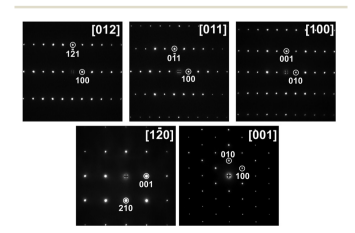


Fig. 2 Electron diffraction patterns for Ta₇Fe₃B₈ along different zone axes.

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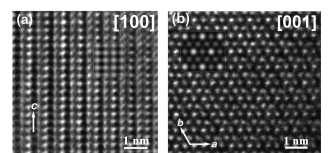


Fig. 3 HRTEM images of $Ta_7Fe_3B_8$: (a) along [100] and (b) along [001] (the two insets within white lines are the simulated images for $\Delta f=-40$ nm and t=6.7 nm along [100] and $\Delta f=-20$ nm and t=6.6 nm along [001], respectively).

powder XRD diffraction (Table 1). There are no superstructure reflections or diffuse reflections characteristic for disorder.

The atomic arrangement in the $TM_7Fe_3B_8$ structure determined from single crystal diffraction (as shown in Fig. 4a) is proved by HRTEM observations (Fig. 3). For the image along [100] (Fig. 3a), the simulated image (the inset in Fig. 3a) was calculated at defocus $\Delta f = -40$ nm and thickness t = 6.7 nm, in which the dark contrast mainly comes from the projection of Ta and Fe atoms. For the HRTEM image along [001] (Fig. 3b; the inset shows the simulated image at a defocus $\Delta f = -20$ nm and thickness t = 6.6 nm), the bright spots are caused by the projection of Ta1 and Fe atoms. HRTEM images also reveal no extended defects in $Ta_7Fe_3B_8$.

Crystal structure

The crystal structures of TM₇Fe₃B₈ adopt a new type of atomic arrangement. As shown in Fig. 4, the structures consist of two alternating layers: the layer of the TM atoms at z = 0 and the layer of the B and Fe atoms at z = 1/2. The B and Fe atoms reside inside [TM₆] trigonal prisms and center [(TM₂)₈] tetragonal prisms, respectively. The trigonal prisms comprise two different types: one is represented by [(B2)(TM1)₂(TM2)₄], and the other one by [(B1)(TM2)₆], respectively. Both represent typical structural building blocks observed in intermetallic borides. The former type of the polyhedra is corrugated and forms hexagonal columns along [001]. Such kind of hexagonal columns has been previously observed in the ternary borides $TM_7TM'_6B_8$ (M = Nb, Ta; M' = Ru, Rh, Ir)⁶² and their disordered variant Ti₇Rh₄Ir₂B₈.⁶³ In TM₇Fe₃B₈, the hexagonal columns are separated by Fe and B1 atoms. This gives rise to the formation of [Fe(TM2)₈] tetragonal prisms and the second type of trigonal prisms. Such buildup of trigonal prims and tetragonal prims in TM₇Fe₃B₈ structures is observed in borides for the first time. This type of structure pattern was previously found in the crystal structure of BaFe2Al9.64 Here the aluminum atoms are occupying the positions of TM2 and Fe, iron atoms are located at the B2 positions, barium is shifted by $[00\frac{1}{2}]$ in respect to Nb1 site, and the positions of B2 are not occupied: Ba₁Al₆Al₃Fe₂□₆ is equivalent to (TM1)₁(TM2)₆Fe₃(B1)₂(B2)₆. Later another ordering variant for

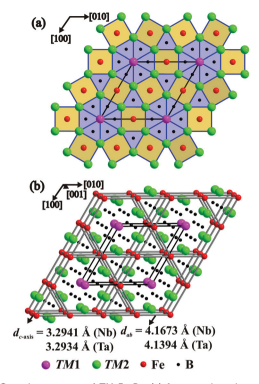


Fig. 4 Crystal structures of $TM_7Fe_3B_8$: (a) framework and packings of polyhedral prisms (pink – TM1 at 1a site, green – TM2 at 6l site, red – Fe at 3g site, black – B1 at 2d site and B2 at 6k site; blue – $[BTM_6]$ trigonal prism, yellow – $[FeTM_8]$ tetragonal prism); (b) three-dimensional Fe-Fe framework and Fe-Fe distances in the (001) plane and along [001].

this atomic motif was discovered in $Hf_5Nb_5Ni_3P_5$: $Hf_1(Hf, Nb)_6(Hf,Nb)_3P_2(Ni,P)_6$.

The structural motif of TM7Fe3B8 can be described as a 2D intergrowth of the AlB₂-type (RX₂ fragment) and CsCl-type (RR' fragment) slabs with the general formula $R_{m+n}R'_mX_{2n}^{66}$ where m = 3 and n = 4 are the numbers of outlined structural fragments per unit cell of TM₇Fe₃B₈. In the AlB₂-type TMB₂ structures, 67,68 condensed [BTM6] trigonal prisms form 3D blocks completely filling the space. Due to such an arrangement, the boron atoms come in close contact and form infinite two dimensional graphite-like nets. Interestingly, in the TM7Fe3B8 structure, the planar six-membered rings of boron are now separated by CsCl-type TMFe slabs, and this also additionally results in the isolated B1 atoms in this structure. Such kind of isolated planar [B₆] rings and the isolated B atoms were also observed in the ternary borides TM7TM'6B862 and their disorder variant Ti₇Rh₄Ir₂B₈.⁶³ Moreover, the TM₇Fe₃B₈ structure is also related to the Mo₂FeB₂-type TM₂FeB₂ structure, 33,34 which can also be described as an intergrowth of the AlB2-type and CsCl-type slabs, now with m = 2 and n = 2 in $R_{m+n}R'_mX_{2n}$. One nearly single-phase Ta₂FeB₂ sample was also synthesized during this study and its structure was refined from powder XRD data.⁶⁹ In TM₂FeB₂, the boron atoms centering the trigonal prism sites come together in B2-pairs, while in TM7Fe3B8, these pairs are fused into [B₆] rings or broken up into isolated B atoms.

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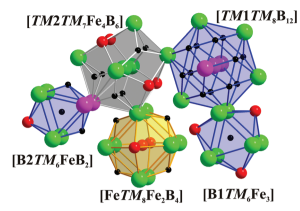


Fig. 5 Coordination polyhedra of the TM, Fe and B atoms in TM₇Fe₇B₈.

Coordination polyhedra of all the atoms in TM₇Fe₃B₈ are depicted in Fig. 5. The polyhedron of TM1 is a 20-vertices polyhedron consisting of 12 B and 8 TM atoms, which could also be observed for TM atoms in TM7TM'6B8,62 TMB2,67,68 and TM₃B₄ ⁷⁰ TM2 and Fe are coordinated by a 17-vertices [TM₇Fe₄B₆] polyhedron and a six-capped [TM₈Fe₂B₄] cube, respectively. The same coordination polyhedra for TM as well as Fe are found in TM₂FeB₂.^{33,34} Finally, B atoms are coordinated by tricapped trigonal prisms formed either by 6 TM and 3 Fe atoms or by 6 TM, 1 Fe and 2 B atoms.

Selected interatomic distances in the crystal structures of TM₇Fe₃B₈ compounds are listed in Table 3. All of them are close or slightly larger than the sum of atomic radii of the elements ($r_{\text{Ta}} = 1.43 \text{ Å}, r_{\text{Nb}} = 1.43 \text{ Å}, r_{\text{Fe}} = 1.24 \text{ Å}, r_{\text{B}} = 0.83 \text{ Å}$). TM-TM contacts in the (001) plane for Nb₇Fe₃B₈ (3.0333(6) Å and 3.0607(4) Å) and Ta₇Fe₃B₈ (3.0053(6) Å and 3.0447(2) Å) are both shorter than these distances in the (001) plane in $TM_7Ru_6B_8$ (3.1257 Å and 3.1189 Å for $Nb_7Ru_6B_8$ and $Ta_7Ru_6B_8$,

Table 3 Selected interatomic distances (Å) in the structures of $Nb_7Fe_3B_8$ and $Ta_7Fe_3B_8$; CN = coordination number

Atoms		Nb ₇ Fe ₃ B ₈	Ta ₇ Fe ₃ B ₈	CN
TM1	-12B2	2.456(3)	2.462(3)	20
	-6TM2	3.0607(4)	3.0447(2)	
	-2TM1	3.2941(1)	3.2934(3)	
TM2	-4B2	2.396(2)	2.383(6)	17
	-2B1	2.4041(3)	2.3921(3)	
	-4Fe	2.7120(2)	2.6995(3)	
	-2TM2	3.0333(6)	3.0053(6)	
	-1TM1	3.0607(4)	3.0447(2)	
	-2TM2	3.0607(4)	3.0447(2)	
	-2TM2	3.2941(1)	3.2934(3)	
Fe	-2B2	2.345(3)	2.310(4)	14
	-2B1	2.4060(1)	2.3899(1)	
	-8TM2	2.7120(2)	2.6995(3)	
	-2Fe	3.2941(1)	3.2934(3)	
B1	-6TM2	2.4041(3)	2.3921(3)	9
	-3Fe	2.4060(1)	2.3899(1)	
B2	-2B2	1.822(5)	1.830(6)	9
	−1Fe	2.345(3)	2.310(4)	
	-4TM2	2.396(2)	2.383(6)	
	-2TM1	2.456(3)	2.462(3)	
		` '	. ,	

respectively)⁶² and AlB₂-type TMB₂ (3.1115 Å and 3.076 Å for NbB₂⁶⁷ and TaB₂,⁶⁸ respectively). TM-TM contacts along the [001] direction for Nb₇Fe₃B₈ (3.2941(1) Å) and Ta₇Fe₃B₈ (3.2934 (3) Å) are both longer than these distances along [001] in TM₇Ru₆B₈ (3.1284(3) Å and 3.1370(3) Å for Nb₇Ru₆B₈ and $Ta_7Ru_6B_8$, respectively)⁶² and TMB_2 (3.2657 Å and 3.275 Å for NbB₂⁶⁷ and TaB₂, ⁶⁸ respectively). B2 atoms in the isolated hexagonal [B₆] rings in TM₇Fe₃B₈ reside slightly off-center in the trigonal prisms towards the Fe atom caps, resulting in longer B2-B2 distances in the hexagonal rings (1.822(5) Å and 1.830(6) Å for Nb₇Fe₃B₈ and Ta₇Fe₃B₈, respectively) than the distances in TMB₂ (1.7962 Å and 1.7759 Å in NbB₂⁶⁷ and TaB₂, 68 respectively), however, still shorter than the distances in the isolated hexagonal rings in $TM_7TM_6'B_8$ (1.868(4) Å, 1.870(6) Å and 1.84(4)-1.87(7) Å in Nb₇Ru₆B₈, Ta₇Ru₆B8 and Nb₇Rh₆B₈).

The iron atoms in the crystal structure of TM₇Fe₃B₈ form a planar Kagome net at z = 1/2. With the relatively large Fe-Fe distances within the Kagome plane (4.17 Å for Nb and 4.14 Å for Ta compound) and the shorter distances of d(Fe-Fe) =3.29 Å along [001] in both compounds (Fig. 4b), this atomic pattern plays important role in the magnetic behavior of these materials (cf. below).

Physical properties

Magnetic properties. The analysis of the magnetic properties of both TM7Fe3B8 compounds is hampered by the presence of ferromagnetic impurities with high Curie temperatures. Fig. 6a shows the magnetic moment μ per formula unit (in Bohr magnetons $\mu_{\rm B}$) for two low fields. Magnetic ordering transitions are clearly visible for both compounds, at $T_N = 240 \text{ K}$ for $Nb_7Fe_3B_8$ and at $T_N = 265$ K for $Ta_7Fe_3B_8$. Below these transitions, tiny remnant magnetization $\leq 0.012\mu_{\rm B}$ for the Nb and $0.006\mu_{\rm B}$ for the Ta compound are also observed. The bulk character of the magnetic transitions is confirmed by sizable anomalies in the specific heat (see below). Further transitions at low temperature are not observed.

Above the ordering transitions, the high-field paramagnetic susceptibilities $\chi(T) = M(T)/H$ were analyzed. First, the intrinsic susceptibility $\chi_{int}(T)$ was extrapolated by the Honda-Owen method from data taken in 3.5 T and 7 T field. Above 250 K, the $\chi_{int}(T)$ data are well described by the Curie-Weiss law (see the plot of H/M in Fig. 6b), $\chi_{int}(T) = C/(T - \theta)$. The effective paramagnetic moment μ_{eff} calculated from C and the Weiss parameter θ are 5.45 $\mu_{\rm B}$ and -195 K for the Nb compound and $5.35\mu_{\rm B}$ and -306 K for the Ta homologue.

Electrical resistivity. In Fig. 7, the electrical resistivities $\rho(T)$ and their temperature derivatives $d\rho/dT$ are shown. Both compounds display metallic conduction with $\rho_{300K} \approx 19 \mu\Omega$ m for the Nb and $\approx 0.55 \mu\Omega$ m for the Ta compound. While the former sample has an extraordinary large resistivity well above the Mott-Ioffe-Regel limit, the latter compound reaches only a value in the range typical for intermetallic compounds. For Nb₇Fe₃B₈ a pronounced kink is visible at the magnetic transition. The kink at T_N is weaker in the Ta compound. The derivatives $d\rho/dT$ indicate that for both compounds a contribution due to spin-disorder scattering of charge carriers is at

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(a) 30
25 Nb 1k0e →
20 TM₇Fe₃B₈
20 15 → 100 0e Nb
1 10 Ta 1k0e →
100 0e Ta
160 160 160 160
180 80 Nb

Fig. 6 (a) Magnetic moment $\mu(T)$ measured in two different fields. The arrows indicate the direction of the temperature sweep. (b) Inverse susceptibility H/M for $\mu_0H=3.5$ T (lower curves for Nb₇Fe₃B₈ and Ta₇Fe₃B₈, respectively) and 7.0 T (middle curves). The data sets above 250 K are the inverse intrinsic susceptibilities $1/\chi_{\rm int}(T)$ (see text). The lines are Curie–Weiss fits to the latter data (see text).

200

7 (K)

400

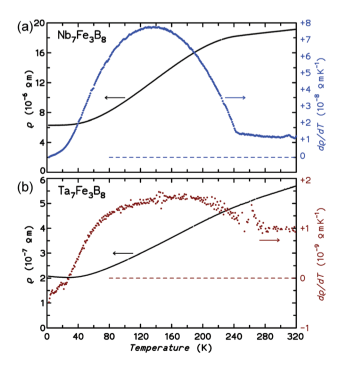


Fig. 7 Electrical resistivity $\rho(T)$ and derivative $d\rho/dT$ for Nb₇Fe₃B₈ (a) and Ta₇Fe₃B₈ (b).

work. Interestingly, the hump in $d\rho/dT$ of the Ta compound extends further to lower temperatures than that of Nb₇Fe₃B₈. The high residual resistivities (RRR value = 3.0 and 2.4 for Nb and Ta-containing compounds, respectively) suggest that both samples have considerable amounts of point defects.

Specific heat capacity. The isobaric specific heat $c_p(T)$ of the $\mathrm{TM}_7\mathrm{Fe}_3\mathrm{B}_8$ compounds is shown in Fig. 8. The strongly bonded light boron atoms in these structures lead to high-frequency optical phonon modes. Therefore, the specific heat at room temperature is still well below the Dulong-Petit limit, *i.e.* $c_p \leq 3nN_\mathrm{A}k_\mathrm{B}$ (n=1) number of atoms in the formula unit, $N_\mathrm{A}=1$ avogadro constant, $N_\mathrm{B}=1$ Boltzmann constant). There are clear second-order anomalies at the weak ferromagnetic ordering transitions. The anomaly for the Ta compound is smaller than that for the Nb homologue. Interestingly, $c_p(T)$ of $\mathrm{Ta}_7\mathrm{Fe}_3\mathrm{B}_8$ is well above that of the Nb compound at temperatures below $\approx 140\,$ K, which may be expected from lower-lying phonon modes of the TM species (Ta has almost twice the atomic mass of Nb).

The inset to Fig. 8 presents the low-T specific heats in a c_p/T vs. T^2 representation. For temperatures below 10 K the $c_n(T)$ may be analyzed following the ansatz $c_p(T) = aT^{-2} + \gamma T + \beta T^3$, where the first term captures the upturn towards the lowest temperatures (observed in the Ta compound only), γT is the contribution from conduction electrons, and βT^3 represents the Debye approximation of the lattice heat capacity. The Sommerfeld coefficients γ are 32.9 and 38.0 mJ mol⁻¹ K⁻¹ for the Nb and Ta compound, respectively. The coefficients β correspond to initial Debye temperatures $\theta_{\rm D}$ of 688 K and 640 K, respectively, the lower $\theta_{\rm D}$ of the latter compound being due to the large atomic mass of tantalum. The origin of the upturn at the lowest T observed for Ta₇Fe₃B₈ is unclear. The application of magnetic fields $\mu_0 H$ of 3, 6, and 9 T leads to a progressive shift of entropy connected to this upturn to higher temperatures, however the involved entropy is very small compared to $N_{\rm A}k_{\rm B}$. This contribution is probably not due to hyperfine split-

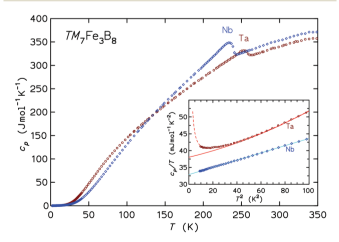


Fig. 8 Specific heat capacity $c_p(T)$. The inset shows the low-T range of the same data in c_p/T vs. T^2 representation. The full lines are fits with electronic and lattice contributions, the dashed line in the case of $Ta_7Fe_3B_8$ also includes an aT^{-2} contribution (see text).

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ting of the nuclear multiplet of 181 Ta (I = 7/2), especially since no corresponding effect is observed for Nb (93 Nb with I = 9/2).

Mössbauer spectroscopy. Spectra of Nb₇Fe₃B₈ at 294 K and 4.3 K are shown in Fig. 9. The Mössbauer spectrum at 294 K shows a single line with small quadrupole splitting (Table 4). Magnetic hyperfine splitting is present at 4.3 K (Fig. 9) with a hyperfine field $H_{\rm hf}$ less than one third of the value for pure α -Fe. The small line width (G/2) (Table 4) indicates the contribution of only one iron species in accordance with the crystal structure. The hyperfine splitting at 4.3 K confirms bulk nature of the magnetic order and suggests that all Fe nuclei experience similar hyperfine fields.

Electronic structure

Both Nb₇Fe₃B₈ and Ta₇Fe₃B₈ are metallic. Their electronic density of states (DOS) features remarkably large contributions of d states of both Nb/Ta and Fe at the Fermi level $(E_{\rm F})$ (Fig. 10). According to the Stoner criterion band splitting and thus spontaneous ferromagnetic ordering are therefore very

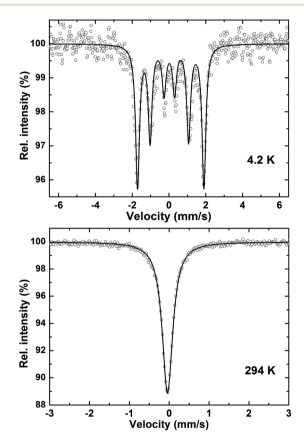


Fig. 9 Mössbauer spectra of Nb₇Fe₃B₈ at 4.3 K and 294 K.

Table 4 Fitting parameters for Mössbauer spectra of Nb₇Fe₃B₈

T(K)	$H_{\rm hf} \left({\rm mm~s}^{-1}\right)$	$eQV_{\rm zz}/4~{\rm (mm~s}^{-1}{\rm)}$	$CS (mm s^{-1})$	$G/2 \text{ (mm s}^{-1}\text{)}$
294	0	-0.020	-0.033	0.158
4.3	3.611	0.038	0.058	0.126

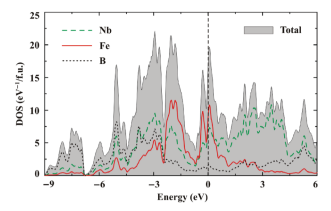


Fig. 10 Spin-unpolarized LDA DOS of Nb₇Fe₃B₈. The Fermi level is at zero enerav.

likely to occur. The Nb/Ta 4d/5d bands are very broad and span the energy range between -5 eV and 5 eV, where clear bonding (Nb and B) and antibonding (Nb only) combinations are formed below and above the Fermi level, respectively. The Fe 3d bands are about half as wide and weakly hybridize with the boron states. The electronic structures of the Nb and Ta compounds are very similar.

The Fermi level is close to a local minimum of the DOS. The number of states at the Fermi level of about 16.5 eV per f.u. and 13.5 eV per f.u. in the spin-unpolarized calculation yields the Sommerfeld coefficients of $\gamma = 38 \text{ mJ mol}^{-1} \text{ K}^{-1}$ and 32 mJ mol⁻¹ K⁻¹ for Nb₇Fe₃B₈ and Ta₇Fe₃B₈, respectively. These values are close to those found experimentally (32.9 and 38.0 mJ mol⁻¹ K⁻¹, respectively), but, surprisingly, reveal a different trend: $\gamma_{Nb} > \gamma_{Ta}$ in DFT, whereas $\gamma_{Nb} < \gamma_{Ta}$ experimentally. Spin fluctuations and magnetic order may affect the number of states at the Fermi level. However, this effect is difficult to assess computationally because the magnetic ground state of TM₇Fe₃B₈ is not known.

Spin-polarized calculations suggest that both Nb₇Fe₃B₈ and Ta₇Fe₃B₈ are magnetic. The magnetic moment on iron is about $1.72\mu_{\rm B}$ regardless of the spin configuration. This value should not be confused with the paramagnetic effective moment of about $3.1\mu_B/Fe$, which is intrinsically higher than the ordered moment and reflects the full fluctuating spin moment, while $1.72\mu_{\rm B}$ revealed by DFT is only its ordered part. On the other hand, this moment is much higher than the tiny remnant magnetization on the order of $0.01\mu_{\rm B}$, which is seen below $T_{\rm N}$.

In order to gain further insight into the nature of the magnetic order in TM7Fe3B8, we analyzed nearest-neighbor exchange couplings by calculating total energies of several spin configurations. The following ordering patterns were considered: ferromagnetic order (I); ferromagnetic order in the ab plane and antiferromagnetic order along c (II); ferrimagnetic (up-up-down) order in the ab plane and ferromagnetic order along c (III). Note that we considered collinear spin configurations only. Therefore, a fully antiferromagnetic order in the ab plane is not possible given the frustrated nature of the kagome spin lattice.

Our spin-polarized calculations revealed the lowest energy of configuration III that we further refer to as zero. The energies of the other two configurations are $E_{\rm I}$ = +65.9 meV per f.u. and $E_{\rm II}$ = +127.1 meV per f.u., respectively. This way, effective nearest-neighbor exchange couplings are J_{ab} = 16.5 meV/Fe and $J_c = -21.2$ meV/Fe, where the positive and negative signs stand for the antiferromagnetic and ferromagnetic couplings, respectively, and we do not divide energies by S^2 because in itinerant magnets it is not a good quantum number. We conclude that the couplings in the ab plane are antiferromagnetic, while the coupling along c is ferromagnetic. Therefore, despite large Fe-Fe distances (cf. above), TM₇Fe₃B₈ are magnetically frustrated, as no collinear spin configuration satisfies the antiferromagnetic couplings in the Kagome net. It is worth noting that effective exchange couplings of +190 K and -247 K are comparable in magnitude to the Curie-Weiss temperature of -195 K in Nb₇Fe₃B₈. However, both ferro- and antiferromagnetic interactions are observed.

Chemical bonding analysis in real space

A striking feature of the TM₇Fe₃B₈ crystal structure is the spatial separation of the TM atoms from the Fe and B ones forming separated planar nets perpendicular to the [001] direction at z = 0 and $z = \frac{1}{2}$ respectively. The reasons for such atomic arrangement were evaluated by the real space analysis of chemical bonding in Nb₇Fe₃B₈ employing electron localizability indicator ELI in its ELI-D representation⁵⁹ (Fig. 11).

While the ELI-D distribution in the penultimate shell of boron atoms has a spherical shape as expected for a p element, the penultimate shells of niobium and iron atoms show strong inhomogeneity being the fingerprint of the participation of these electrons in the bonding interactions in the valence region. 60,72 In the valence region around B1 atoms reveal five ELI-D maxima (attractors). Three of them are located on the Fe-B contacts visualizing the 2c(Fe-B) bonds.

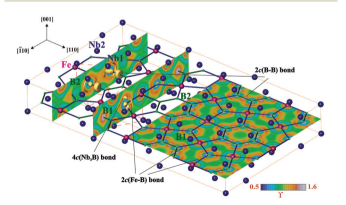


Fig. 11 Electron localizability indicator in Nb₇Fe₃B₈: (left and middle) ELI-D (Υ) distribution in the planes perpendicular to the Fe-B net revealing the structuring of the penultimate shells of Nb1, Nb2 and Fe atoms as well as location of the ELI-D attractors (shown by the yellow isosurface with $\Upsilon = 1.315$) around the boron atoms within the [BNb₆] trigonal prisms (red dashed line); (right) ELI-D distribution in the plane at $z = \frac{1}{2}$ with the maxima visualizing the two-center B-B and Fe-B bonds.

The basins of the remaining two attractors are located above and below the boron nucleus being in contact with the core basins of three Nb atoms beside ones of B1. This arrangement reflects the four-center interaction. ELI-D reveals similar distribution around B2. According to the local symmetry, two of the attractors in the plane at $z = \frac{1}{2}$ visualize the 2c(B-B) bonds, the third one shows mainly 2c(Fe-B) interaction. The basins of the attractors above and below the plane are not present, being united with that of the Fe-B interaction indicating here a delocalization of a 2c bond toward a multicenter one. Thus, the plane of Fe and B atoms at $z = \frac{1}{2}$ with its Kagome topology mentioned above is formed mainly by two-center interactions. Between this plane and the niobium atoms at z = 0 the multicenter bonding is observed. Assuming that the multicenter bonding is weaker than the two-center interactions, such bonding picture should yield a pronounced cleavage of the material perpendicular to the [001] direction.

Conclusions

In this study, two new ternary borides $TM_7Fe_3B_8$ (TM = Nb, Ta) with Kagome-type iron sublattices were synthesized by arcmelting of the elements and subsequent annealing at 1500 °C. Their hexagonal primitive structure is an intergrowth of AlB₂type and CsCl-type slabs, involving [BTM₆] trigonal prisms and [FeTM₈] tetragonal prisms. The condensation of trigonal prisms results in the formation of hexagonal columns along c-axis, hence, also forming isolated planar [B₆] rings in this structure.

Metallic character of TM7Fe3B8 is confirmed by temperature dependence of the electrical resistivity as well as by the sizable linear term in the specific heat for both compounds.

Magnetic susceptibility measurements reveal predominantly antiferromagnetic order below $T_N = 240 \text{ K}$ for Nb₇Fe₃B₈ and T_N = 265 K for Ta₇Fe₃B₈. The sextet in the Mössbauer spectra of $Nb_7Fe_3B_8$, the sizable anomalies in the specific heat at T_N , and the kinks in the resistivity curves confirm the bulk character of the magnetic transitions for both compounds. These transitions are related to the presence of sizable magnetic moments localized on the Fe atoms within the planar Kagometype iron sublattice. DFT calculations indicate metallic behaviour for both compounds and show that interactions in the ab plane are antiferromagnetic and thus subject to a strong geometrical frustration that should prevent Néel type magnetic ordering. On the other hand, strong interplane coupling (of any sign) can effectively suppress this frustration and trigger the formation of long-range-ordered states,⁷³ which is probably the case in TM₇Fe₃B₈. However, the presence of remnant magnetization unanticipated in a regular Kagome antiferromagnet indicates a more complex nature of the magnetic order.

Quantum-chemical analysis of the chemical bonding in Nb₇Fe₃B₈ within the electron localizability approach reveals five ELI-D maxima around B1, visualizing three in-layer Fe–B bonds and two interactions with core basins of three Nb atoms above and below, while only three maxima around B1,

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showing two B-B bonds and one Fe-B interaction. The analysis also indicates that the Kagome net of Fe and B is mainly formed by two-center interactions, whereas multicenter bonding between this net and Nb atoms is observed.

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References

- 1 V. I. Matkovich, Boron and Refractory Borides, Springer, New York, 1977.
- 2 K. Upadhya, J. M. Yang and W. P. Hoffman, Am. Ceram. Soc. Bull., 1997, 76, 51-56.
- 3 J. B. Levine, S. H. Tolbert and R. B. Kaner, Adv. Funct. Mater., 2009, 19, 3519-3533.
- 4 R. G. Munro, J. Res. Natl. Inst. Stand. Technol., 2000, 105, 709-720.
- 5 P. Schwarzkopf, R. Kieffer, W. Leszynski and K. Benesovsky, Refractory Hard Metals, Borides, Carbides, Nitrides, and Silicides, Macmillan, New York, 1953.
- 6 C. Buzea and T. Yamashita, Supercond. Sci. Technol., 2001, 14, R115-R146.
- 7 W. G. Fahrenholtz, G. E. Hilmas, I. G. Talmy and J. A. Zaykoski, J. Am. Ceram. Soc., 2007, 90, 1347-1364.
- 8 A. Inoue, Acta Mater., 2000, 48, 279-306.
- 9 J. M. Lafferty, J. Appl. Phys., 1951, 22, 299-309.
- 10 J. F. Herbst, Rev. Mod. Phys., 1991, 63, 819-898.
- 11 K. Moorjani and J. M. D. Coey, Magnetic Glasses, Elsevier, Amsterdam, 1984.
- 12 C. H. Smith, ed. in *Rapidly* solidified alloys, H. H. Liebermann, Marcel Dekker, New York, 1993.
- 13 L. Pauling, The Nature of the Chemical Bond and the Structure of Molecules and Crystals: An Introduction to Modern Structural Chemistry, Cornell University Press, Ithaca, 3rd edn, 1960.
- 14 B. Albert and H. Hillebrecht, Angew. Chem., Int. Ed., 2009, 48,8640-8668.
- 15 T. Mori, in Handbook on the Physics and Chemistry of Rare Earths, ed. K. A. Gschneidner, J.-C. Bünzli and V. Pecharsky, North-Holland, Amsterdam, 2008, vol. 38, pp. 105–173.
- 16 P. Rogl and H. Nowotny, J. Less-Common Met., 1978, 61, 39-

- 17 H. Nowotny and P. Rogl, in Boron and Refractory Borides, ed. V. Matkovich, Springer, New York, 1977, ch. 23, pp. 413-438.
- 18 Yu. B. Kuz'ma, Crystallochemistry of Borides, Lviv University Publishers, Lviv, 1983.
- 19 P. Rogl, in *Inorganic Reactions and Methods*, ed. A. P. Hagen, Wiley, 1991.
- 20 Q. Zheng, F. R. Wagner, A. Ormeci, Yu. Prots, U. Burkhardt, M. Schmidt, W. Schnelle, Yu. Grin and A. Leithe-Jasper, Chem. - Eur. J., 2015, 21, 16532-16540.
- 21 M. Ade and H. Hillebrecht, Inorg. Chem., 2015, 54, 6122-6135.
- 22 J. M. Vandenberg, B. T. Matthias, E. Corenzwit and H. Barz, Mater. Res. Bull., 1975, 10, 889-894.
- 23 H. C. Ku, G. P. Meisner, F. Acker and D. C. Johnston, Solid State Commun., 1980, 35, 91-96.
- 24 B. T. Matthias, E. Corenzwit, J. M. Vandenberg and H. E. Barz, Proc. Natl. Acad. Sci. U. S. A., 1977, 74, 1334-1335.
- 25 Q. Zheng, R. Gumeniuk, H. Rosner, W. Schnelle, Yu. Prots, U. Burkhardt, Yu. Grin and A. Leithe-Jasper, J. Phys.: Condens. Matter, 2015, 27, 415701.
- 26 Q. Zheng, W. Schnelle, Yu. Prots, M. Bobnar, U. Burkhardt, A. Leithe-Jasper and R. Gumeniuk, Dalton Trans., 2016, 45, 3943-3948.
- 27 D. Gignoux and D. Schmitt, in Handbook of Magnetic Materials, ed. K. H. J. Buschow, North-Holland, Amsterdam, 1997, vol. 10, p. 239.
- 28 R. Dronskowski, K. Korczak, H. Lueken and W. Jung, Angew. Chem., Int. Ed., 2002, 41, 2528-2532.
- 29 O. Isnard and D. Fruchart, J. Alloys Compd., 1994, 205, 1-15.
- 30 B. P. T. Fokwa, Eur. J. Inorg. Chem., 2010, 2010, 3075-
- 31 M. Mbarki, R. St. Touzani and B. P. T. Fokwa, Angew. Chem., Int. Ed., 2014, 53, 13174-13177.
- 32 Yu. B. Kuz'ma, Dopov. Akad. Nauk URSR, Ser. A, 1967, 16, 939-940.
- 33 Yu. B. Kuz'ma, T. I. Ts'olkovskii and O. P. Baburova, Inorg. Mater. (Engl. Trans.), 1968, 4, 950-953.
- 34 Yu. B. Kuz'ma, A. S. Sobolev and T. F. Fedorov, Sov. Powder Metall. Met. Ceram., 1971, 10, 410-414.
- 35 P. Rogl, in Iron Systems: Phase Diagrams, Crystallographic and Thermodynamic Data, ed. G. Effenberg and S. Ilyenko, Springer, 2008, vol. 11D1.
- 36 L. Balents, Nature, 2010, 464, 199-208.
- 37 S. Yan, D. A. Huse and S. R. White, Science, 2011, 332, 1173-1176.
- 38 O. Janson, J. Richter and H. Rosner, Phys. Rev. Lett., 2008, 101, 106403.
- 39 M. P. Shores, E. A. Nytko, B. M. Bartlett and D. G. Nocera, J. Am. Chem. Soc., 2005, 127, 13462-13463.
- 40 A. Dönni, G. Ehlers, H. Maletta, P. Fischer, H. Kitazawa and M. Zolliker, J. Phys.: Condens. Matter, 1996, 8, 11213-11229.
- 41 P. Mendels and F. Bert, J. Phys. Soc. Jpn., 2010, 79.

Paper

42 T.-H. Han, J. S. Helton, S. Chu, D. G. Nocera,

- J. A. Rodriguez-Rivera, C. Broholm and Y. S. Lee, Nature, 2012, 492, 406-410.
- 43 M. Fu, T. Imai, T.-H. Han and Y. S. Lee, Science, 2015, 350, 655-658.
- 44 H. Ishikawa, Y. Okamoto and Z. Hiroi, J. Phys. Soc. Ipn., 2013, 82, 063710.
- 45 I. Rousochatzakis, J. Richter, R. Zinke and A. A. Tsirlin, Phys. Rev. B: Condens. Matter, 2015, 91, 024416.
- 46 WinXPow, STOE and Cie GmbH, Darmstadt, 2003.
- 47 L. Akselrud and Yu. Grin, J. Appl. Crystallogr., 2014, 47, 803-805.
- 48 More information about NbFeB and TaFeB single crystals are given in ESI.† The metallography analysis, crystal structure and magnetic data for NbFeB and TaFeB single crystals are shown in Fig. S1 to Fig. S3,† respectively. Refined crystallographic data from powder XRD patterns of NbFeB and TaFeB crytals are given in Tables S2 and S3.†
- 49 J. L. Pouchou and F. Pichoir, in Electron Probe Quantitation, ed. K. F. J. Heinrich and D. E. Newbury, Plenum Press, New York, 1991, ch. 31-70.
- 50 V. Grillo, STEM-Cell program.
- 51 S. Mørup and E. Both, in Nuclear Instruments and Methods, Elsevier, Amsterdam, 1975, pp. 445-448.
- 52 K. Koepernik and H. Eschrig, Phys. Rev. B: Condens. Matter, 1999, **59**, 1743–1757.
- 53 J. P. Perdew and Y. Wang, Phys. Rev. B: Condens. Matter, 1992, 45, 13244-13249.
- 54 O. Jepsen, A. Burkhardt and O. K. Andersen, The Program TB-LMTO-ASA, version 4.7, Max-Planck-Institut für Festkörperforschung, Stuttgart, 1999.
- 55 U. von Barth and L. J. Hedin, J. Phys. C: Solid State Phys., 1972, 5, 1629-1641.

- 56 O. K. Andersen, Phys. Rev. B: Solid State, 1975, 12, 3060-3083.
- 57 W. R. L. Lambrecht and O. K. Andersen, Phys. Rev. B: Condens. Matter, 1986, 34, 2439-2449.
- 58 M. Kohout, Int. J. Quantum Chem., 2004, 97, 651-658.
- 59 M. Kohout, Faraday Discuss., 2007, 135, 43-54.
- 60 F. R. Wagner, V. Bezugly, M. Kohout and Y. Grin, Chem. -Eur. I., 2007, 13, 5724-5741.
- 61 M. Kohout, DGrid, version4.6, 2011.
- 62 Q. Zheng, M. Kohout, R. Gumeniuk, N. Abramchuk, H. Borrmann, Yu. Prots, U. Burkhardt, W. Schnelle, L. Akselrud, H. Gu, A. Leithe-Jasper and Yu. Grin, Inorg. Chem., 2012, 51, 7472-7483.
- 63 B. P. T. Fokwa and M. Hermus, Angew. Chem., Int. Ed., 2012, 51, 1702-1705.
- 64 K. Turban and H. Schäfer, J. Less-Common Met., 1975, 40, 91-96.
- 65 H. Kleinke and H. F. Franzen, J. Am. Chem. Soc., 1997, 119, 12824-12830.
- 66 E. I. Gladyshevsky and Yu. Grin, Sov. Phys. Crystallogr., 1981, 26, 683-689.
- 67 S. Okada, K. Hamano, T. Lundström and I. Higashi, AIP Conf. Proc., 1991, 231, 456-459.
- 68 S. Okada, K. Kudou, I. Higashi and T. Lundström, J. Cryst. Growth, 1993, 128, 1120-1124.
- 69 a = 5.8619(2) Å, c = 3.2762(2) Å; Ta at (4 h) (0.1797, x + 1/2, 1/2), Fe at (2a) (0, 0, 0), and B at (4f) (0.3880, x + 1/2, 0).
- 70 R. Kiessling, Acta Chem. Scand., 1949, 3, 603-615.
- 71 J. Emsley, The Elements, Clarendon Press, Oxford, 1998.
- 72 M. Kohout, R. F. Wagner and Yu. Grin, Theor. Chem. Acc., 2002, 108, 150-156.
- 73 O. Götze and J. Richter2015, arXiv:1510.04898.