# Interplay between crystal structure and magnetism in Tb2CoGe2 upon hydrogenation

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The intermetallic compound Tb2CoGe2 adopts the structure type Sc2CoSi2 (Pearson symbol *mS*20, space group *C*2/*m*) [1], which belongs to the intergrowth structure series based on CrB- and TiNiSi-type slabs. The representatives of this structure series can be described by the general formula *Rm*+*nTmMm*+*n* (*R* is a rare-earth metal, *T* is a *d*-element, *M* is a main group element, *m* and *n* represent the number of TiNiSi and CrB fragments, respectively). In the case of Sc2CoSi2-type compounds the ratio *m* : *n* = 2 : 2 [2] (Fig. 1a).

The initial intermetallic Tb2CoGe2 was synthesized by arc-melting the constituent metals. Hydrogenation of the activated (by heating in vacuum) crushed sample was performed under a hydrogen pressure of 770 mbar. Heating up to 250°C was required to initiate the reaction. The pressure drop in the system corresponded to the hydride composition Tb2CoGe2H0.9. Phase analysis and crystal structure determinations were performed by X-ray powder diffraction (Bruker D8 Advance diffractometer, Cu *K*α radiation). A SQUID magnetometer (Quantum Design) was used for magnetic measurements in the temperature range 2-300 K and in magnetic fields of up to 7 T.

The refined crystal structure parameters of Tb2CoGe2 are close to those reported in [1]: space group *C*2/*m*, *a* = 10.5276(5) Å, *b* = 4.2062(4) Å, *c* = 10.1888(5) Å, *β* = 118.083(2)°, *V* = 398.05(5) Å3, Tb1 (4*i*) (0.0078(5) 0 0.3329(4)), Tb2 (4*i*) (0.1888(4) 0 0.1105(4)), Co (4*i*) (0.2641(8) 0 0.6425(14)), Ge1 (4*i*) (0.3464(8) 0 0.4339(8)), Ge2 (4*i*) (0.4785(8) 0 0.1171(8)). The structure of the metallic matrix is preserved upon hydrogenation: *C*2/*m*, *a* = 10.9455(5) Å, *b* = 4.1356(2) Å, *c* = 10.1555(5) Å, *β* = 119.189(2)°, *V* = 401.33(4) Å3, Tb1 (4*i*) (-0.0058(5) 0 0.3197(4)), Tb2 (4*i*) (0.1893(5) 0 0.0968(5)), Co (4*i*) (0.2761(11) 0 0.6192(12)), Ge1 (4*i*) (0.3493(8) 0 0.4285(8)), Ge2 (4*i*) (0.5064(9) 0 0.1361(8)). Hydrogenation results in a moderate cell volume increase (∆*V*/*V* = 0.82 %) caused by lattice expansion in the direction of the *a*-axis (∆*a*/*a* = 3.97 %), partly compensated by lattice contraction in the directions of the *b*- (∆*b*/*b* = -1.68 %) and *c*-axes (∆*c*/*c* = -0.33 %).

Tb2CoGe2 exhibits multiple magnetic phase transitions below the ordering temperature *T* = 130 K (Fig. 1b,c). Based on the shape of the magnetization curve at 2 K, the ground state can be assumed to be ferrimagnetic, while the magnetization curve at 150 K is typical for an approach to antiferromagnetism. Hydrogenation suppresses magnetic interactions in the hydride, which orders magnetically at 25 K. The results will be discussed in terms of different impact of hydrogenation on the environment of Tb atoms in different positions.



###### **Figure 1.** Representation of the crystal structure of Tb2CoGe2 as intergrowth of CrB- and TiNiSi-type slabs (a); magnetic susceptibility curves for Tb2CoGe2 and Tb2CoGe2H0.9 measured in field-cooled (FC) and zero-field-cooled (ZFC) mode (*b*); magnetization curves for Tb2CoGe2 and Tb2CoGe2H0.9 measured at various temperatures (*c*).

#### [1] Bodak, O. I., Pecharskii, V. K., Starodub, P. K., Salamakha, P. S., Mrooz, O. Y. & Bruskov, V. A. (1986). *Izv. Akad. Nauk SSSR, Met.* **4**, 216.

#### [2] Zhao, J. & Parthé, E. (1989). *Acta Crystallogr.* *C* **45**, 1853.

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