Structure and hard magnetic properties of TbCu7-type SmFe8.95−xGa0.26Nb x nitrides

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Nanocrystalline SmFe8.95−xGa0.26Nb x (x = 0, 0.1, 0.2, 0.3) were prepared using rapid-quenching, annealing and nitriding. The magnetic properties and crystal structures were systematically studied under various wheel velocities to investigate the influence of Nb doping for the compounds. It is found that TbCu7-type structure is able to be obtained even though the wheel velocity is reduced to 20 m/s (x = 0.3). An significant increase (∆T c = 70 °C) of the Curie temperature is obtained with Nb doping at x = 0.1 due to the lattice expansion revealed by Rietveld analysis. The optimum coercivity with the value H c0 of 810 kA/m is achieved at x = 0.2 in the nitrides, in which a reasonable distribution of grain sizes of both TbCu7-type SmFe9N2 and α-Fe can be found. However, an excess of Nb doping may lead to the increase of the weight fraction of α-Fe, which in turn deteriorates the magnetic properties. © 2018 Published by Elsevier B.V. on behalf of Chinese Society of Rare Earths.

1. Introduction

Since Coey et al.1 discovered intermetallic compounds Sm2Fe7N14, a systematical and profound study about them has been conducted. About one year later, Katter et al.2 first discovered hexagonal TbCu7-type phase in melt spin ribbons of binary Sm−Fe system. Stoichiometry of the TbCu7-type phase is about SmFe9, which has moderate coercivity upto 2.13 kOe. However, the TbCu7-type phase is a meta-stable phase, and only at high cooling speed it can be obtained. In order to enhance and stabilize the magnetic properties, material researchers have already made some successful attempts by substitution of 3d and boron elements for iron.3−7 The additional 3d element can facilitate TbCu7-type structure and improve magnetic properties, such as V, Ti, Zr, Al.8−11 Recently, our group12 reported that the moderate doping content of Ga for SmFe9N2 compounds with TbCu7-type structure was very effective to increase Curie temperature T c and coercivity. However, the problem is that obtaining a single TbCu7 phase still requires high velocity (50 m/s) and the magnetic property is relatively low.

This work is proposed to reduce the wheel velocity of obtaining a single TbCu7 structure by adding Nb and further improve the magnetic properties as well based on the results mentioned above. Single phase materials of SmFe8.95−xGa0.26Nb x were prepared, and the effects of Nb addition on the crystal structure, Curie temperature and hard magnetic properties were studied in the Nb doping TbCu7-type Sm−Fe−Ga nitrides.

2. Experimental

SmFe8.95−xGa0.26Nb x (x = 0, 0.1, 0.2, 0.3) parent alloys were prepared by induction melting Sm (99.9%), Fe (99.9%), Ga (99.9%) and Nb (99.9%) in Ar gas atmosphere. An extra amount of 25 wt% Sm was added to compensate the weight losses during melt-spinning and induction melting process. The molten ingots were ejected through an orifice of 0.7 mm in diameter at the bottom of a quartz crucible. All ribbons were prepared by melt spinning onto a rotating Cu-disk at a wheel speed of velocities 20−50 m/s (intervals 10 m/s). Subsequently, the ribbons wrapped in tantalum-foil were annealed at 800 °C for 1 h in vacuum (<1.0 × 10−3 Pa) and rapidly quenched with room temperature water. Then the annealed SmFe8.95−xGa0.26Nb x powders were nitrided at 450 °C for 16 h in nitrogen atmosphere. The crystal structures and phases in the specimens were examined by X-ray diffraction (XRD) using Co Kα radiation by continuous (2θ = 0.02°, 20°−90°) and step mode (2θ = 0.02°, 20°−110°, counting time 4 s, 9 kW). The data were...
processed by Rietveld refinement with Rigaku Plus software. The thermomagnetic curves of the samples were tested using a vibrating sample magnetometer (VSM, Quantum Design VersaLab) with an applied field of 1000 Oe. Magnetic powder properties were measured at room temperature using a VSM in fields up to 3T. Microstructures of as-annealed ribbons were observed by transmission electron microscopy (TEM).

3. Results and discussion

3.1. Structure and phase analysis

Fig. 1 shows some X-ray diffraction patterns for SmFe$_{8.95-x}$Co$_x$Ga$_{0.26}$Nb$_x$ ($x = 0, 0.1, 0.2, 0.3$) alloys quenched at various wheel velocities of 20–50 m/s (interval = 10). It has been shown that the magnetic phases are obvious changed, depending on the Nb content and the speed of rapid-quenching velocities. For SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ($x = 0$) alloy, the single TbCu$_7$-type structure was able to be obtained when the speed of velocities reach to 50 m/s. However, when the speed of velocities is lower than 50 m/s, the main phase is the rhombohedral Th$_2$Zn$_{17}$-type structure with a small amount of SmFe$_2$ and a-Fe phases which is from Fig. 1(a). With the content of Nb increasing, the lowest speed of velocities to obtain single TbCu$_7$-type structure gradually decreased as shown in Fig. 1. Furthermore, the single TbCu$_7$-type structure was able to be obtained even though the speed of velocities was 20 m/s with Nb doping at $x = 0.3$, which can be seen from Fig. 1(d). The reason can be attributed to the increasing undercooling of SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ alloys by adding high melting point metal Nb, which can impel the formation of the TbCu$_7$-type structure in the melt spun Sm–Fe alloys. However, an excess of Nb doping may lead to the increased content of $\alpha$-Fe and precipitation of the Nb$_2$Fe$_7$ phase with main TbCu$_7$-type phase, which in turn deteriorates the magnetic properties, which is in agreement with the literature that the independence of formation of the TbCu$_7$-type phase on the wheel velocities in the range of 15–50 m/s for the melt spun Sm$_{10}$Fe$_{82.5}$V$_{7.5}$ alloys, due to the substitution effect of vanadium for iron, can impel the formation of the TbCu$_7$-type structure in the melt spun Sm–Fe alloys. As a result, we concluded that the magnetic phases are sensitively dependent both on the composition and the wheel velocity and the TbCu$_7$-type phase shows a tendency to form in higher wheel speed, which is different from the situation of the Th$_2$Zn$_{17}$-type structure.$^{15}$

The crystal structure of the as-annealed ribbons has been refined by Rietveld method based on the TbCu$_7$-type structure. The observed patterns agree well with the final calculated patterns. The results are shown in Table 1, which illustrates the variation of lattice parameters and axial ratio $c/a$ value with the content of Nb element ($v = 40$ m/s). With the increase of Nb doping ($x \leq 0.2$), the lattice parameter $a$ decreases and the $c$ increases, as a result, a greater expansion of $c/a$ value is observed. In general, the main phase of TbCu$_7$-type structure tends to form in the compounds when the $c/a$ value dropped in the range of 0.8499–0.8699, which is in agreement with the that of Katter et al.$^2$ And the greater the ratio of $c/a$ is, the more stable the TbCu$_7$-type crystal structure will be.$^7,16$ On the contrary, an excessive doping of Nb ($x = 0.3$) can lead to a decline in $c/a$, corresponding to a trend of instability for TbCu$_7$ structure.

![Fig. 1. X-ray diffraction patterns of as-annealed SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ (x = 0 (a), 0.1 (b), 0.2 (c), 0.3 (d)) ribbons prepared at various wheel velocities of 20–50 m/s.](image-url)
Fig. 2 shows thermomagnetic behaviors for as-annealed ribbons in a magnetic field of 1000 Oe (v = 40 m/s). Remarkably, the Curie temperature of the as-annealed SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ribbons increases from 470 K for $x = 0$–540 K for $x = 0.1$. To certify it, the average iron moment $<\mu_{Fe}>$ for each compound is calculated at the dumbbell sites from the saturated moments, based on the hypothesis that an opposing magnetic moment of each samarium is equal to 0.90 $\mu_B$. It can be found that $<\mu_{Fe}>$ increases slightly from 1.66 $\mu_B$ for $x = 0$ to 1.72 $\mu_B$ with Nb doping of $x = 0.1$ shown in Fig. 3. It can be attributed to the enhanced Fe–Fe interaction, which is consistent with increase of the Curie temperature. Furthermore, the Curie temperatures of 540 K for SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ($x = 0.1$) is much higher than that of SmFe$_{12}$ or Sm$_5$Fe$_{17}$ based alloys previously reported.

3.2. Nitrogenation and magnetic properties

Nitrides of SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ($x = 0, 0.1, 0.2, 0.3$) alloys were obtained by quenching at 40 m/s, crystallization annealing at 800 °C for 60 min in vacuum ($<1.0 \times 10^{-3}$ Pa) and subsequent nitriding at 450 °C for 16 h in nitrogen atmosphere. Fig. 4 shows the hysteresis loop of the nitride powders of SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ($x = 0, 0.1, 0.2, 0.3$), which were measured in an maximum applied field of ±30 kOe. The greatest magnetic property of $B_{r} = 0.76$ T, $H_{cj} = 810$ kA/m, and $(BH)_{max} = 80$ kJ/m$^3$ was obtained in the nitride powders of SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ribbons at $x = 0.2$. Magnetic energy product has been improved by 32 kJ/m$^3$ and the intrinsic coercivity increased by 320 kA/m compared to Nb-free. It can be seen that the demagnetization curve “step” become smaller, which means the increase of magnetic performance. This can be explained from two aspects. On the one hand, the maximum $c/a$ ratio is obtained for $x = 0.2$, which can improve the efficiency of nitriding, and the maximum $c/a$ ratio increases interstitial lattice of 3f site occupied by N atoms, according to Teresiak et al. Fig. 5(a) shows XRD patterns of SmFe$_{8.75}$Ga$_{0.26}$Nb$_{0.2}$ alloys with different conditions treatment in order to investigate the structure after nitriding. The diffraction peaks of as-annealed Sm–Fe alloys are similar to that of as-quenched sample. But the diffraction peaks of the nitrides shift toward the left because N atoms penetrate into the gap of crystal lattice. Fig. 5(b) shows the thermomagnetic behaviors for as-annealed and nitridation of SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ alloys in a magnetic field of 1000 Oe. The Curie temperature of the as-annealed SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_{0.2}$ ribbons increases from 420 K for $x = 0$–750 K after nitriding.

On the other hand, the microstructure of the as-annealed SmFe$_{8.75}$Ga$_{0.26}$Nb$_{0.2}$ ribbons is more homogeneous than that of Nb-free ribbons according to Fig. 6(b). Furthermore, we can see the grain size of nitridation SmFe$_{8.95-x}$Ga$_{0.26}$Nb$_x$ ($x = 0.2$) has not changed in Fig. 6(c). The grain size of SmFe$_{8.75}$Ga$_{0.26}$Nb$_{0.2}$ ribbons is about 35 nm, much smaller than the grain size (62 nm) of Nb-free alloys.
magnetization application and subsequent removal of direct doping of Nb (\(x = 0.3\)) shows up a steep step, which results in the deterioration of the remanence as well as the coercivity, the reason is that excess \(x\) can result in a slight decline in \(\alpha\) and the increased contents of \(x\)-Fe as indicated from XRD Rietveld refinement results displayed in Table 2, which means a trend of instability for TbCu7 structure.

A small amount of \(x\)-Fe emerges for each SmFe8.95-xGa0.26Nb_x compound as displayed in Table 2. Henkel plots for the \(x = 0.2\) sample are selected to investigate intergranular exchange coupling existing in the SmFe8.95-xGa0.26Nb_xN_y compounds. The expression which is the interaction between soft magnetic phase and hard magnetic phase was given as follows23:

\[
\delta_m(H) = \frac{\left[M_H(H) - M_r + 2M_r(H)\right]}{M_r}
\]

Where the remanent magnetization \(M_r(H)\) is acquired after the application and subsequent removal of direct field \(H\), the remanent magnetization \(M_H(H)\) after DC saturation in one direction and the subsequent application and removal of a direct field \(H\) in the reverse direction. Fig. 6 shows the Henkel plots of the SmFe8.75-Ga0.26 Nb_x powders. Obviously, there is no exchange coupling indicated by the negative peak of \(\delta_m(H)\) which is observed in Fig. 7. The non-existing exchange coupling may be ascribed to the relatively larger grain size of \(x\)-Fe phase in the present work due to the relatively lower wheel velocity of 40 m/s. The grain size of \(x\)-Fe phase determined by the XRd Rietveld in Table 2 is approximately in nm, which is somewhat difficult to expect the hard magnetic properties by the exchange coupling phenomena.24 It is in agreement with the report by Katter et al.8 that the exchange coupling existing in the nanocomposite magnetic system was obtained in the higher wheel velocity of 60 m/s.

4. Conclusions

The crystal structures and hard magnetic properties of melt-spun SmFe8.95-xGa0.26Nb_x ribbons and their nitrides were studied to illustrate the influence of Nb substitution for Fe in the compound. Firstly, the single TbCu7-type structure is able to be obtained at the relatively low velocity of 20 m/s because of Nb doping \((x = 0.3)\). Second, an increase \((\Delta T_m = 70\degree C)\) of the Curie temperature is obtained with Nb doping at \(x = 0.1\) compared to the Nb-free samples. Thirdly, the optimum coercivity with the value \(H_c\) of 810 kA/m is gained at \(x = 0.2\) in the nitrides. Finally, an excess of Nb doping will deteriorate the magnetic properties instead, which may be ascribed to the increase of the weight fraction of \(x\)-Fe.

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**Fig. 6.** TEM micrographs of as-annealed SmFe8.95Ga0.26 (a), micrographs of as-annealed SmFe8.95Ga0.26Nb0.2 (b) and TEM micrographs of SmFe8.95Ga0.26Nb0.2N. (c).

**Fig. 7.** The Henkel plots of the SmFe8.95Ga0.26Nb0.2N powders.

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**Table 2**

| Composition | Fe grain size/nm | Content Fe wt% |
|-------------|-----------------|---------------|
| x = 0       | 50.1            | 7.3           |
| x = 0.1     | 31.7            | 1.3           |
| x = 0.2     | 30              | 0.72          |
| x = 0.3     | 28.1            | 2.5           |

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The table shows the grain size of Fe phase in as-annealed SmFe8.95-Ga0.26 alloy obtained from the Rietveld refinement.
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