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Magnetic Properties and Magnetic Domain Observation of Nd–Fe–B Sintered Magnets Treated by Grain Boundary Diffusion Process with Dy–Al Co-Sorption

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Effective grain boundary diffusion (GBD) process with Dy–Al co-sorption is applied to enhance the coercivity of Nd–Fe–B sintered magnets. The coercivity of the magnet ($H_{cJ} = 1789 \text{ kA} \cdot \text{m}^{-1}$) subjected to the present GBD treatment was observed to be superior to that of the untreated magnet ($H_{cJ} = 1003 \text{ kA} \cdot \text{m}^{-1}$) and the conventional GBD magnet ($H_{cJ} = 1661 \text{ kA} \cdot \text{m}^{-1}$) treated with DyAl alloy. In the present GBD magnet, the Dy–Al co-sorption process facilitated Dy diffusion into the center region of the magnet (thickness: 3.5 mm), resulting in high coercivity. Further, magnetic domain observations were made using magnetic force microscopy (MFM) to observe the thermal demagnetization behavior of the present GBD magnet. The present GBD magnet suppressed the continuous domain reversal of adjacent grains; thus, the partially persistent single-domain structure remained, even at 453 K. [doi:10.2320/matertrans.MT-M2020389]

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1. Introduction

The addition of heavy rare-earth (HRE = Dy or Tb) elements in Nd–Fe–B sintered magnets efficiently enhances the coercivity $(H_{cJ})^{1-3}$ but reduces the maximum energy product, $(BH)_{Max}$, because of the reduction in the remanence (B_r) due to the antiferromagnetic coupling between the HRE elements and Fe in the (Nd,Dy)₂Fe₁₄B main phase. Recently, the price of these sintered magnets has been increasing because of insufficient HRE resources, which is a major problem in the industrial field. Furthermore, given the increasing awareness of environmental issues, magnets with high coercivity, particularly those applicable in electric vehicles (EV/HEV), are in high demand.⁴⁾ Therefore, it is necessary to improve the coercivity of Nd–Fe–B sintered magnets while minimizing the amount of HRE used.

The process of selectively enriching the HRE elements in the grain boundary region surrounding the Nd₂Fe₁₄B main phase, that is, the core-shell type microstructure, is an efficient way to reduce the amount of HRE in the Nd–Fe–B sintered magnets. To obtain a suitable core-shell type microstructure, a grain boundary diffusion (GBD) process in which the magnets were coated by HRE elements and subsequently heated above the melting point of the Nd-rich grain boundary was developed.^{5–18)} Following heat treatment, the resulting magnets exhibit high coercivity and suppress the decrease in remanence.

We previously reported a GBD process in which Dy or Tb metals were coated on the magnet surface by a sputtering method⁷⁾ or a reduction diffusion method applied to HRE compounds and Ca reductants.^{8,17)} Using these techniques, a structure in which the Nd₂Fe₁₄B main phase is surrounded by the HRE-enriched grain boundary was achieved. Consequently, the coercivity was effectively enhanced without a severe reduction in the remanence. We also reported

improved magnetic properties of Nd-Fe-B scrap powder by Yb metal sorption (both adsorption and absorption), utilizing the low boiling point of the HRE elements.⁹⁾ Similar GBD processes using various HRE compound powders,¹⁰⁻¹³⁾ vapor deposition, 5,6,14-16) and alloy powder^{18,19} have also been reported. In particular, a GBD process using Dy-Ni-Al alloy powder was reported by Oono et al.;18) the simultaneous Dy diffusion with Ni and Al yielded a decrease in the melting point of the grain boundary. As a result, Dy diffusion into the deeper region of the magnets was facilitated, resulting in high coercivity, even with a thickness of 5 mm. However, the preparation of the alloy powder requires largescale equipment for melting and grinding the alloy, and the fine alloy powder containing HRE elements is extremely active in air atmosphere; thus, handling and mass productivity become a problem.

Meanwhile, in the GBD process, the HRE elements diffuse from the magnet surface, causing a difference in the HRE concentration near the surface and deep regions in the magnets, which limits both the applicable magnet size and the increase in coercivity. In mass production, to achieve high coercivity, the GBD process with Tb raw material, which has a higher magnetic anisotropy and is more expensive than Dy, is widely applied.

Referring to a previous study,¹⁸ to obtain a more effective coercivity enhancement of Nd–Fe–B sintered magnets using the GBD process with Dy, which is cheaper than Tb, this study focuses on the melting point of the grain boundary and investigates the simple GBD process with Dy–Al co-sorption. Dy (b.p. = 2835 K^{20}) and Al (b.p. = 2740 K^{21}) are low-boiling-point elements, and sorption readily occurs under reduced pressure conditions such as $1.0 \times 10^{-3} \text{ Pa}.^{22}$ Conventionally, Al has been used as a minor additive element in Nd–Fe–B sintered magnets.^{23,24} The diffusion of Al into the Nd-rich grain boundary yields a decrease in the melting point,²⁵ and Dy diffusion into the deeper region of the magnet is expected.

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Although there are numerous reports on the grain boundary microstructure of GBD magnets,^{5,6,14,15)} there are several reports on the thermal demagnetization behavior at high temperatures, which are important properties for the driving motors of electric vehicles. Therefore, to investigate the demagnetization behavior in more detail, magnetic domain observations using magnetic force microscopy (MFM) were also performed to observe the thermal demagnetization behavior of the present GBD magnet treated by Dy–Al co-sorption.

2. Experimental

In this study, the conventional and present GBD processes were applied. The conventional GBD process applied Dy sorption, DyAl alloy, and a mixture of TbF₃ and CaH₂. The procedure for Dy sorption and a mixture of TbF3 and CaH2 were based on previous reports.^{8,16)} The DyAl alloy with a molar ratio of Dy:Al = 1:1 was prepared by arc melting, grinding to approximately 5 µm, then coating onto the magnet surface via anhydrous-butanol. The present GBD process applied Dy-Al co-sorption. Figure 1 shows a typical experimental procedure for the present GBD process. Massproduced Nd–Fe–B sintered magnets ($B_r = 1.43$ T, $H_{cJ} =$ 1003 kA·m⁻¹) were cut into square-shaped $(3.0 \times 3.0 \times$ 2.8 mm³) or cylindrical $(10^{\phi} \times 3.5^{t} \text{ mm}^{3})$ pieces by machining. The oxide layer on the magnet surface was subsequently removed by etching with dilute nitric acid for several minutes. Al metal powder (99.0% purity) with an average particle diameter of approximately 30 µm and anhydrousbutanol were mixed at a weight ratio of 50:50. After immersing the magnets in this slurry solution, hot air drying was performed to uniformly coat the Al powder on the magnet surface. The Al-coated magnet was vacuum-sealed with Dy metal grains (purity 99.9%) in a quartz tube maintained at a vacuum degree of 1.0×10^{-3} Pa. The quartz tubes were heated at 1173-1273 K for 6 hrs and subsequently annealed at 823 K for 2 hrs. After heat treatment, the re-



Fig. 1 Typical experimental procedure for the present GBD process.

sulting GBD magnets were ultrasonically washed with dilute nitric acid and ethanol to remove impurities on the magnet surface.

The magnetic properties were evaluated using a pulseexcitation-type BH (PBH) tracer or a superconducting magnet-based vibrating sample magnetometer (SCM-VSM) with a magnetic field range of $\pm 4775 \text{ kA} \cdot \text{m}^{-1}$. The SCM-VSM measurement method was described in previous papers.^{26,27)} Microstructural characterizations were performed using an X-ray diffractometer (XRD), field emission electron probe micro-analyzer (FE-EPMA), scanning transmission electron microscope (STEM). The chemical composition was quantitatively analyzed by inductively coupled plasma (ICP) optical emission spectrometry. Magnetic domain observations were performed using MFM. A high coercivity probe with a coercivity of $>796 \text{ kA} \cdot \text{m}^{-1}$ was applied. By following the probe based on the pre-scanned shape images, the distance between the magnet surface and the probe can remain constant at approximately 20-30 nm. The MFM observations were performed in situ after increasing the temperature from 300 K to 373 K and 453 K in a vacuum degree of approximately 1.0×10^{-4} Pa and maintaining it for 1.5 hrs at each predetermined temperature. Before the observation, the magnets were magnetized by a pulsed magnetic field of 6366 kA·m⁻¹ after flat-mill-polishing on the magnet surface. Note that, since the evaluations of the magnetic properties and the MFM observations were performed at different site facilities, the magnetized magnets were vacuum-packed to enable transportation.

3. Results and Discussion

3.1 XRD patterns and magnetic properties

Figure 2 shows the XRD patterns measured on the magnet surface (a) treated by the present GBD process with Dy–Al co-sorption at 1223 K for 6 hrs and (b) untreated. Here, the untreated magnet referred to a magnet that was sintered and subsequently annealed. Note that, acid washing or polishing was not performed before XRD. In the present GBD magnet, the peak intensity of the fcc NdO phase (measured lattice constant: a = 0.5025 nm) was clearly increased compared to



Fig. 2 XRD patterns of the magnet surface (a) treated by the present GBD process with Dy–Al co-sorption at 1223 K for 6 hrs and (b) untreated.



Fig. 3 Dependence of magnetic properties $(H_{c,J}, B_r, \text{ and } (BH)_{Max})$ on the heat treatment temperature for the present GBD magnet.

the untreated magnet. This indicated that an Nd-rich phase was regenerated on the magnet surface by the GBD process. However, the peak intensity of the fcc NdO phase was extremely weak in the untreated magnet, that is, the Nd₂Fe₁₄B main phase was almost bare on the magnet surface before the GBD process. This was presumably due to the oxidation of the Nd-rich phase caused by machining or surface oxidation, and subsequently peeled off the magnet surface. Meanwhile, the regeneration of the Nd-rich phase was due to Dy, which was vapor-deposited on the magnet surface and diffused into the inner region by substituting with Nd, which was enriched in the grain boundary. This is similar to the case where Nd was sputter-deposited on the magnet surface and heat treatment was subsequently applied.²⁸⁾ This result indicated that, like Nd, Dy recovered the magnet surface, which deteriorates owing to oxidation and machining. While, Al powders pre-coated on the magnet surface also became vapor during heat treatment under the present conditions. However, because Al powders were coated directly on the magnet surface, Al sorption proceeded preferentially compared to Dy sorption.

Figure 3 shows the dependence of magnetic properties $(H_{cJ}, B_r, \text{ and } (BH)_{Max})$ on the heat treatment temperature for the $3.0 \times 3.0 \times 2.8 \text{ mm}^3$ magnets treated by the present GBD process. The heat time was constant at 6 hrs, and the values were evaluated by the PBH tracer. The coercivity (H_{cJ}) increased with increasing heat treatment temperature, and the remanence (B_r) and maximum energy product $((BH)_{Max})$ decreased because of the reduction of the saturation magnetization (M_s) by proceeding with the interstitial diffusion of Dy into the Nd₂Fe₁₄B main phase. In particular, when the heat treatment temperature increased to 1273 K, which is close to the melting point of the Nd₂Fe₁₄B main phase, the coercivity became highly enhanced from $1003 \text{ kA} \cdot \text{m}^{-1}$ to $1835 \text{ kA} \cdot \text{m}^{-1}$. In addition, even at a heat temperature of 1173 K, the coercivity ($H_{cJ} = 1571 \text{ kA} \cdot \text{m}^{-1}$) exceeded the conventional GBD process applying Dy as a diffusion material, such as mixed powder of $DyF_3 + CaH_2$ $(H_{cJ} = 1420 \text{ kA} \cdot \text{m}^{-1}).^{8)}$ Based on the results, we can select the heat treatment conditions according to the required coercivity and remanence. In this study, the basic heat



Fig. 4 Demagnetization curves of the magnets measured by SCM-VSM (a) untreated, treated by the conventional GBD process with (b) Dy sorption,
(c) DyAl alloy, (d) mixture of TbF₃ and CaH₂, and (e) treated by the present GBD process with Dy–Al co-sorption. Heat treatment conditions remain constant at 1223 K for 6 hrs.

Table 1 Summary of the magnetic properties of the (a) untreated magnet, (b)–(d) conventional GBD magnets, and (e) the present GBD magnet.

	H_{cJ}	B_r	H_k/H_{cJ}	$(BH)_{Max}$
	/ kA•m ⁻¹	/ T	(%)	/ kJ•m ⁻³
(a) untreated	1003	1.43	91	388
(b) Dy sorption	1641	1.41	77	378
(c) DyAl alloy	1661	1.39	89	375
(d) TbF_3 and CaH_2	1761	1.40	90	387
(e) Dy-Al co-sorption	1789	1.38	85	342

treatment temperature was set to 1223 K based on the increased coercivity and decreased ratio of remanence.

Figure 4 shows the demagnetization curves of the untreated and treated (conventional and present GBD processes) magnets evaluated by SCM-VSM. The heat treatment conditions were constant at 1223 K for 6 hrs and subsequently annealed at 823 K for 2 hrs. H_{cl} , B_r , H_k/H_{cl} , and $(BH)_{Max}$ are summarized in Table 1. Here, H_k is the value of the magnetic field required to reduce the remanence (B_r) by 10%. The magnets were cylindrical, with dimensions of $10^{\phi} \times 3.5^{t}$ mm³. Independent of the details of the GBD process, the coercivity was enhanced while suppressing the reduced remanence. This demonstrated that GBD process is a suitable process for manufacturing high-coercivity magnets. The present GBD magnet exhibited the highest coercivity among the studied magnets. This increase in coercivity is the same as that of the conventional GBD process with a mixed powder of TbF₃ and CaH₂; that is, the present GBD process is effective for the substitution of Tb by Dy. Interestingly, the increase in coercivity was higher than that of the DyAl alloy containing Al as a diffusion material, as well as Dy-Al cosorption. The reasons for this are discussed in the microstructure section. On the other hand, the GBD process with uniformly coated powders on the magnet surface with the (c) DyAl alloy or (d) TbF₃ and CaH₂ yielded a higher squareness



Fig. 5 BSE images and EPMA images of Nd, Dy, and Al at various depths from the magnet surface (a) untreated, (b) treated by the conventional GBD process with DyAl alloy and (c) treated by the present GBD process with Dy–Al co-sorption. (d) Dy line profile of each GBD magnet obtained by EPMA line analysis.

 (H_k/H_{cJ}) than the GBD process including Dy sorption (b) and (e). This is because of the uniform coating of HRE-containing powders on the magnet surface, which enabled more uniform diffusion of HRE into the magnets.

3.2 Observation of grain boundary microstructure

Figure 5 shows the backscattered electron (BSE) images and EPMA images of Nd, Dy, and Al at various depths from the untreated and treated (conventional GBD process with DyAl alloy and present GBD process with Dy–Al co-sorption) magnet surfaces, and the Dy line profile of each GBD magnet obtained by EPMA line analysis. The compositions of Nd, Dy, and Al for each magnet quantitatively analyzed by ICP are summarized in Table 2. In each GBD magnet, Dy was excessively enriched around the Nd₂Fe₁₄B main phase near the magnet surface, particularly in the case of the conventional GBD process with DyAl alloy, the Dy diffusion into the main phase was remarkable (see Fig. 5(b) and (c) 0 mm-Dy). On the other hand, in the central region of the magnet, Dy diffusion was

Table 2 Composition of Nd, Dy, and Al of the magnet (a) untreated, treated by (b) the conventional GBD process with DyAl alloy and (c) the present GBD process with Dy–Al co-sorption, measured by ICP.

	Composition / wt.%		
	Nd	Dy	Al
(a) untreated	31.22	< 0.01	0.15
(b) DyAl alloy	30.45	1.17	0.39
(c) Dy-Al co-sorption	30.37	1.20	0.61

clearly observed in the present GBD process with Dy–Al co-sorption, but was slight in the DyAl alloy (see Fig. 5(b) and (c) 1.75 mm-Dy). Although we have reported various GBD processes,^{7,8,17)} the diffusion distance from the magnet surface was several tens of micrometers to several hundreds of micrometers when Dy was applied as a diffusion material. Therefore, notably, the present GBD process exhibited sufficient Dy diffusion up to 1.75 mm from the magnet

surface. This is also confirmed by Fig. 5(d), which shows the diffusion distance of Dy from the magnet surface. In the DyAl alloy, the Dy diffusion was pronounced near the surface to approximately 0.1 mm, but the amount of Dy reaching the center region was higher in the Dy-Al co-sorption than in the DyAl alloy. Therefore, the diffusivity of Dy in the present GBD process is superior to that of the conventional GBD process. Lu et al.¹⁹ reported that the GBD process with Tb-Pr-Cu-Al alloy applied to the magnet surface improves the wettability and mobility of the diffusion alloy during heat treatment owing to the effect of Al, resulting in high coercivity. In addition, they also suggested that the Al content of the diffusion alloy reached 25 at%, Tb and Al diffusion became more pronounced near the magnet surface where it was in direct contact with the diffusion alloy. This is in good agreement with the results of the DyAl alloy containing 50 at% Al in this study, that is, a large amount of Dy was consumed by excessive Dy diffusion into the Nd₂Fe₁₄B main phase near the magnet surface, which prevents sufficient Dy diffusion into the center region of the magnet. On the other hand, in the present GBD process, Al was not alloyed with Dy and diffused preferentially compared to Dy. Therefore, the microstructure or chemical properties of the grain boundary is expected to change, but the wettability or mobility of the diffusion material itself should not be directly affected as in the case of DyAl alloy. Thus, in the present GBD process with Dy-Al co-sorption, a deeper Dy diffusion into the center region of the magnet was achieved, resulting in an effective enhancement of the coercivity. In the present GBD magnet, it was confirmed that, in addition to Dy, Al diffused along the grain boundary to the center region of the magnet compared to the conventional GBD magnet (see Fig. 5(b) and (c) 1.75 mm-Al).

Then, STEM observations were applied to investigate the grain boundary microstructure in more detail. STEM observations were performed around the center region of the magnets. Figure 6 shows STEM high-angle annular-darkfield (HAADF) images of the magnet (a) untreated and (b) treated by the present GBD process with Dy-Al co-sorption. The grain boundary of the untreated magnet was amorphous and thin, but the present GBD magnet was thicker than the untreated magnet and showed a periodic pattern, suggesting the existence of a crystalline phase. The thicknesses of the grain boundary before and after the GBD process were approximately 1.4 and 2.4 nm, respectively. The reason for the thicker grain boundary of the present GBD magnet is that Nd was substituted by diffused Dy near the surface of the Nd₂Fe₁₄B main phase and precipitated Nd was enriched in the grain boundary, as previously reported.^{13,14} Figures 6(c)-(e) shows the fast Fourier transform (FFT) patterns taken from the grain boundary of the present GBD magnet, and each pattern was obtained from the region indicated by the white circles in Fig. 6(b). The FFT patterns show a discrete ring pattern (denoted by yellow arrows in Fig. 6(c)-(e)), suggesting the existence of an amorphous phase in the grain boundary. Note that, since a beam diameter applied to the FFT was narrowed down to 2 nm, the ring pattern showing an amorphous phase was discrete. This ring pattern was also slightly observed, even at the grain boundary, indicating a symmetric lattice fringe in the STEM image (see Fig. 6(d)).

The FFT patterns in Fig. 6(d) also show a non-ferromagnetic fcc NdO [1 1 2] phase with a measured d-space of the (1 1 -1) plane for 0.293 nm (0.288 nm for the International Center for Diffraction Data (ICDD): Card No. 00-033-0937). STEM observations revealed that the grain boundary of the present GBD magnet comprised an fcc NdO phase and an amorphous phase. To date, many researchers have observed the microstructure of the Nd₂Fe₁₄B grain boundary to obtain guidelines for increasing the coercivity. Commonly reported that the grain boundary contains Nd-rich amorphous phase,^{14,15,29,30)} several Nd oxide phases,^{31,32)} and Cu-rich phase.^{14,15,29,30)} Watanabe *et al.*^{5,6)} applied the GBD process with Tb-vapor sorption and reported that the resulting Ndrich phase consisted mainly of a crystalline NdO₂ phase and a minor amorphous phase including Fe, Co, and Cu, which is in good agreement with the results of this study. Thus, there are various reports on the grain boundary microstructure, and further research is required. Figure 6(f) shows the STEM-HAADF image and STEM-EDX elemental mappings for Nd, Dy, Al, Fe, Cu, Co, and O of the present GBD magnet. The elemental mappings show that Dy, Al, Cu, and Co were also enriched in the Nd-rich grain boundary, while Fe was diluted. The FFT patterns demonstrated the existence of an amorphous phase and an fcc NdO phase in the grain boundary, and STEM-EDX elemental mapping revealed the enrichment of Dy, Al, Cu, and Co, similar to Nd. Combining these results, Dy, Al, Cu, and Co in the grain boundary existed as an amorphous phase.

Figure 7 shows the STEM-HAADF image of the present GBD magnet and the concentration line profile of Fe, Nd, O, and Al. Co, Cu, and Dy near and at the grain boundary were extracted by STEM-EDX area-analysis. The STEM-EDX was collected in the yellow box shown in Fig. 7(a). The concentration shown in Fig. 7(a) was integrated along the vertical line in the box. Note that, Fig. 6(f) and Fig. 7(a) are for the same region. Compared with the Nd₂Fe₁₄B main phase, the concentration of Fe in the grain boundary decreased, while the other elements increased, and the concentration of Nd was 24.9 at%. For Dy and Al which applied as a diffusion material in the present GBD process, the concentrations increased to 1.3 at% and 3.4 at%, respectively. While the thickness of the grain boundary of the present GBD magnet was approximately 2.4 nm (see Fig. 6(b)), the concentration profile shows that the full width at half maximum (FWHM) of Dy and Al was about 4 nm. This indicates that Dy and Al were enriched beyond the grain boundary to near the surface of the Nd₂Fe₁₄B main phase, which enhances the coercivity but decreases the remanence, as mentioned above. According to a previous study,²⁾ substituting Nd with Dy in the Nd₂Fe₁₄B main phase enhances the magnetic anisotropy field (H_A) from 5332 $kA \cdot m^{-1}$ to 11937 $kA \cdot m^{-1}$, but decreases the saturation magnetization (M_s) from 1.60 T to 0.71 T. Assuming that the (Nd,Dy)₂Fe₁₄B shell comprises Nd and Dy in the ratio of 24.9:1.3 at%, its magnetic anisotropy is estimated at 5660 kA \cdot m⁻¹, which is an increase of 6.1%, and its saturation magnetization is estimated at 1.56 T, which is a decrease of 2.5%. Note that, this estimation is based on the assumption that the composition of the $Nd_2Fe_{14}B$ main phase core is Dy-free Nd₂Fe₁₄B. However, the actual increase ratio in the



Fig. 6 STEM high-angle-annular-dark-field (HAADF) images of the magnet (a) untreated and (b) treated by the present GBD process with Dy–Al co-sorption. (c)–(e) Fast Fourier transform (FFT) patterns taken from the grain boundary of the present GBD magnet. Each pattern is obtained from the region indicated by the white circles in (b) and (f) STEM-HAADF image and STEM-EDX elemental mappings for Nd, Dy, Al, Fe, Cu, Co, and O of the present GBD magnet.

coercivity was much higher than the estimated increase ratio in the magnetic anisotropy field. This may be attributed to the difference in Dy concentration between the surface and center regions of the magnet, particularly because the center region where the estimation was applied, the Dy concentration was lower compared to the surface region. In addition, it is also considered that there are other factors, such as changing the grain boundary microstructure, which contributes to the increase in coercivity. Meanwhile, the actual decrease ratio in the remanence was slightly higher than the estimated decrease ratio in the saturation magnetization. The saturation magnetization reportedly decreased due to the substitution of Fe by Al at the j_2 site of the Nd₂Fe₁₄B phase,³³⁾ this may be one reason for the present result.

Then, we discuss the role of Al, which is an important element in this study. Oono *et al.*¹⁸⁾ suggested that the existence of diffused elements such as Ni and Al in the grain boundary may contribute to the deep diffusion of Dy by

decreasing the melting point of the grain boundary. Combining the experimental results, that is, microstructure observation and concentration profiling, with this speculation, the melting point of the grain boundary is expected to decrease by approximately 430 K in the phase diagram of the Nd–Al binary system.²⁵⁾ As a marked decrease in the melting point of the grain boundary, the diffusivity of Dy improved during the GBD process by actively transforming the grain boundary into the liquid-state grain boundary. Hence, a deeper Dy diffusion was achieved, and the coercivity was effectively enhanced.

3.3 Thermal demagnetization and magnetic domain observation

Figure 8 shows the coercivity measured at 300 K, 373 K, and 453 K for the magnet (a) treated by the present GBD process with Dy–Al co-sorption and (b) untreated, evaluated by SCM-VSM. The coercivity of the present GBD magnet at



Fig. 7 STEM-HAADF image of the present GBD magnet (a) and concentration line profile of (b) Fe, Nd, O and (c) Al, Co, Cu, and Dy near and the grain boundary extracted by STEM-EDX area-analysis. The STEM-EDX was collected in the yellow box shown in (a) and the concentration was integrated along the vertical line in the box.



Fig. 8 Coercivity measured at 300 K, 373 K, and 453 K for the magnet (a) treated by the present GBD process with Dy–Al co-sorption and (b) untreated.

373 K was similar to that of the untreated magnet at 300 K. In addition, the temperature coefficient of coercivity β also improved from -0.53%·K⁻¹ to -0.50%·K⁻¹, indicating that the present GBD magnet improved the heat resistance in accordance with the increase in coercivity.

To study the behavior of thermal demagnetization in more detail, magnetic domain observations using MFM were performed. The MFM observations were performed on a polished surface perpendicular to the *c*-axis, and the polishing depth from the surface was approximately 100 μ m. In these observation regions, the magnetic anisotropy field was effectively enhanced by the GBD process. The MFM images for the untreated magnet at each observation temperature are shown in Fig. 9(a) and schemati-



Fig. 9 MFM images and schematic drawings obtained at 300 K, 373 K and 453 K for the thermal demagnetized magnet (a), (b) untreated and (c), (d) treated by the present GBD process with Dy–Al co-sorption.

Table 3 Demagnetization ratio of the magnet (a) untreated and (b) treated by the present GBD process with Dy–Al co-sorption at each temperature.

Demagnetization ratio (%)				
Temperature / K	(a) untreated	(b) Dy-Al co-sorption		
300	47.4	11.1		
373	60.3	13.6		
453	89.7	61.7		

cally drawn in Fig. 9(b). In the schematic, the magnetic domain state of each grain is shown in each color. The MFM images and schematic drawings of the present GBD magnet obtained by the same method are shown in Fig. 9(c) and 9(d). The demagnetization ratios at each observed temperature are summarized in Table 3. The demagnetization ratio is defined as follows:

Demagnetization ratio (%)

$$=$$
 the number of multi-domain grains
the number of total grains – Nd-rich phase × 100

At 300 K, despite being magnetized at $6366 \text{ kA} \cdot \text{m}^{-1}$, a significant number of multi-domain grains with maze patterns were observed. The existence of such multiple domains in the initial state may be due to the existence of grains with almost zero coercivity, which are caused by a local decrease in the

magnetic anisotropy field by polishing the magnet surface, as reported by Kobayashi et al.³⁴⁾ The demagnetization ratios at 300 K were 47.4% and 11.1% in the untreated and present GBD magnets. In the untreated magnet, the adjacent grains appeared to have a continuous multi-domain structure, whereas in the present GBD magnet, the multi-domain structure did not exhibit such strong continuity; rather, they appear to be isolated. This may be because the nonferromagnetic fcc NdO phase in the present GBD magnet existed in the grain boundary, which contributed to the magnetic decoupling between the Nd₂Fe₁₄B main phases, and also enhanced the magnetic anisotropy field by enriching Dy at the surface of the Nd₂Fe₁₄B main phase. The surface defects of the Nd₂Fe₁₄B main phase, that is, magnetically coupled grains, and the high local demagnetization stray field at the sharp corners and edges of grains, are reportedly the cause of the decrease in the magnetic anisotropy field.^{35–38)} In the present GBD magnet, such surface defects in the Nd₂Fe₁₄B main phase were repaired by the formation of a thick grain boundary and a high magnetic anisotropy field $(Nd,Dy)_2Fe_{14}B$ shells, thereby suppressing magnetization reversal. Bance et al.³⁹⁾ reported that (Dy,Nd)₂Fe₁₄B shells with a thickness of 4 nm around the Nd₂Fe₁₄B main phase canceled out the reduction of the coercivity owing to the existence of surface defects. The thickness of these shells are in good agreement with the approximately 4 nm FWHM of the Dy concentration profile confirmed by the present GBD magnet. (see Fig. 7(c)) In fact, in the present GBD magnet with high coercivity, most of the magnetic domains persisted the single-domain state at 373 K, but the untreated magnet demagnetized by 60.3%. Finally, at 453 K, 89.7% of the untreated magnet and 61.7% of the present GBD magnets were found to have a multi-domain structure due to thermal demagnetization. Although these results show a similar trend to the thermal degradation of coercivity at the same temperature shown in Fig. 8, there was a difference of approximately 10% in the absolute value of the demagnetization ratio. This may be because the comparison is based on the total coercivity measured as the magnets and the results of the magnetic domain observations on a local surface region that is susceptible to demagnetization.

4. Summary

We applied an effective GBD process with Dy-Al cosorption to Nd-Fe-B sintered magnets. The coercivity of the present GBD magnet increased from $1003 \text{ kA} \cdot \text{m}^{-1}$ to $1789 \text{ kA} \cdot \text{m}^{-1}$. This increase in coercivity is higher than that of conventional GBD magnets treated with DyAl alloy $(H_{cJ} = 1661 \text{ kA} \cdot \text{m}^{-1})$ and $\text{TbF}_3 + \text{CaH}_2$ $(H_{cJ} = 1761 \text{ kA} \cdot \text{m}^{-1})$ $kA \cdot m^{-1}$). The present GBD process facilitated Dy diffusion into the center region of the magnet with a thickness of 3.5 mm, resulting in high coercivity. This is presumably because the preferential Al diffusion decreased the melting point of the grain boundary, thereby improving the diffusivity of Dy. STEM observations revealed that Dy and Al were clearly enriched in the grain boundary, even in the center region of the magnet. Furthermore, it was also demonstrated that the grain boundary comprised of an fcc NdO phase and an amorphous phase including Dy, Al, Cu, and Co. As an extension study, the thermal demagnetization behavior was observed using MFM. In the present GBD magnet, the existence of a non-ferromagnetic fcc NdO phase, which contributes to magnetic decoupling between the Nd₂Fe₁₄B main phases, and the formation of (Dy,Nd)₂Fe₁₄B shells with a high magnetic anisotropy field suppressed the continuous domain reversal of adjacent grains. As a result, in the present GBD magnet, a partially persistent single-domain structure was observed, even at 453 K.

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