PROCESSING Sm-Fe(Ta)-N HARD MAGNETIC MATERIALS

PROCESIRANJE TRDIH MAGNETOV NA OSNOVI Sm-Fe(Ta)-N

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SmFe based al loys in tersti tially mod i fied with ni trogen are potential can di dates for high en ergy per manent magnets. In or der to ob tain the op ti mum prop er ties a thor ough un der stand ing of the start ing ma ter rial and pro cess ing parameters is required. The micro struc tures of two cast al loys of com po si tion Sm_{138} Fe $_{822}$ Ta₄₀ and Sm_{137} Fe $_{863}$ were carefully ex am ined with a SEM equipped with EDX and the ex act stoichiometries of the phases were de ter mined. The SmFeTa ma terial was found to con tain sig nif i cant amounts of TaFe 2, as well as the Sm₂ Fe $_{17}$, SmFe 2, SmFe 3 phases ob served in the SmFe ma terial but with out the sig nif i cant amounts of TaFe 2, as well as the Sm₂Fe₁₇, SmFe 2, SmFe 3 phases ob served in the SmFe ma te rial but with out the α -iron dendrites which are char ac ter is tic of the lat ter ma te rial. Hy dro gen ab sorp tion-de sorption stud ies car ried out on both ma ter i als have dem on strated the in creased re sis tance to de com position of the Sm₂(FeTa) $_{17}$ material, requiring an additional 150°C for the re ac tion to go to com ple tion. The op ti mum con di tions neces sary to provide the high est coercivities us ing the con ventional HDDR process com bined with pre-milling were in ves ti gated. The coercivities ob tained af ter us ing the HDDR process and sub se quent nitriding with out any pre-milling were 680 kA/m for the SmFeTaN and 360 kA/m for the SmFeN sam ples. Sig nif i cantly higher coercivites of 1000 kA/m for SmFeN and 1275 kA/m for SmFeTaN were achieved by re duc ing the part ic le size with mill ing prior to the HDDR process. The better coercivities ob tained with the Ta con tain ing sam ple were found to be due to the pres ence of a much smaller amount of α Fe. The mill ing prior to the HDDR treat ment im proves the coercivity be cause of the small part ic le size, which pre vents the grains grow ing too large, with their con sequent very neg a tive effect on the coercivity. ef fect on the coercivity.

Key words: SmFeN, Ta, HDDR, mill ing

Zlitine na osnovi sistema Sm-Fe, intersticijsko modificirane z du{ikom, imajo velik potencial kot trajni magnetni materiali. Optimalne magnetne lastnosti lah ko dose emo s pravilno obdelavo zlitine, ki pa temelji na poznavanju materiala. Primerjali smo zlitini sestave Sm₁₃₈ Fe₈₂₂ Ta₄₀ in Sm₁₃₇ Fe₈₆₃. Fazno sestavo smo preu-evali z elektronsko vrsti~no mikroskopijo in EDX analizo. SmFeTa zlitina vsebuje TaFe 2 fazo, poleg nje pa tudi Sm₂Fe ₁₇, SmFe 2, SmFe 3 faze, ki so prisotne tudi v SmFe zlitini z izjemo, da SmFe zlitina namesto TaFe 2 vsebuje dendritsko αFe. [tudirali smo procese absorbcije in desorbcije vodika. Ugotovili smo, da Ta stabilizira Sm₂Fe ₁₇ fazo, saj je bila temperatura razpada te faze za 150°C vi{ja od tem per a ture, pri kateri je razpadla Sm₂Fe ₁₇ faza iz binarne zlitine. Poiskali smo optimalne pogoje postopka HDDR, ki so nam dali naj bolj{e re zultate na podro-ju magnetnih lastnosti. Koercitivnosti prahov dobljenih po tem postopku in nadaljnjem nitrira nju, so 680 kA/m za Sm₁₃₈ Fe ₈₆₃ N_x. Ob-utno vi{je koercitvnosti smo dosegli z mletjem materiala, ki smo ga izvedli nred nostonkom HDDR. t.i. 1275 kA/m za Sm₁₃₈ Fe ₈₆₂ Ta₄₀ N_x in 1000 kA/m za Sm₁₃₈ Fe ₈₆₃ N_x. koercitivnosti pred postopkom HDDR, t.j. 1275 kA/m za Sm₁₃₈ Fe $_{82,2}$ Ta₄₀ N_x in 1000 kA/m za Sm₁₃₈ Fe $_{86,3}$ N_x. Razliko v vi{ji koercitivnosti sestave s Ta, gre pripisati manj{i vsebnosti α -Fe. Vi{je koercitivnosti pred mletega materiala pa dejstvu, da v majhnih delcih rast velikih zrn, ki imajo negativen vpliv na koercitivnost, ni mo`na.

Klju~ne besede: SmFeN, Ta, HDDR, mletje

1INTRODUCTION

SmFeN magnets have received considerable attention since their discovery in 1991¹ Their intrinsic properties are comparable with, or better than, those of magnets based on Nd₂Fe₁₄B, and so they have the potential to take a significant share of the rare earth permanent magnet market. SmFeN magnets are open to a number of possible processing routes, for example, mechanical alloying^{2,3}, melt spinning^{4,5} and conventional powder metallurgy^{1,6} however the HDDR process⁷⁸ appears to be the most promising.

The addition of a third element, in this case Ta, is primarily intended to reduce the incidence of α -iron which typically constitutes some 25% of the phases present⁹ in the as-cast binary alloy. The α -iron can be removed by an extended vacuum or inert gas heat treatment, however this is expensive, environmentally detrimental and can cause problems with maintaining precise composition as a result of samarium evaporation. The addition of 4-5% Nb¹⁰ or Ta¹¹ or 1% Zr^{12} to the

procedure.

SmFe material allows the primary crystallisation of α -iron to be avoided.

Effects of reducing the particle size in combination with the HDDR process have been noted by other workers, Müller et al ¹³ found that by milling after the HDDR process was completed a remarkable increase in coercivity could be observed. This increase was attributed to the well known effects of size reduction as a result of milling. They also found milling prior to the HDDR process had a minimal effect unless an additional 5 wt% of samarium was added. In this case the improvement was attributed to a more even microstructure not possible with larger particles due to having grain formation begin earlier in the centre of the large particles. Gebel et al ¹² also used vibration milling under toluene to produce fine powders prior to applying the HDDR process as part of their experimental

In this paper we have investigated the effects of a Ta addition on the hydrogen absorption behaviour as a function of temperature using Thermopiesic Analysis (TPA) and Gas Flow Analysis (GFA). We have also produced magnets using powders with a variety of pre-milling times combined with a conventional HDDR and nitriding process to assess the benefits of the Ta addition.

2EXPERIMENTAL

Both the SmFe and SmFeTa cast alloys were produced by conventional induction melting methods in 5 kg batches by Less-Common Metals Ltd. Small pieces of approximately 0.5 cm³ were cut from similar parts of each alloy and mounted and polished for metallographic examination. The results of this examination have been published previously¹⁴ and the analyses on the Sm₂Fe₁₇ phase showed that 2-3% of Ta is dissolved in this phase. As cast materials were also investigated using x-ray diffractometry to determine their lattice parameters.

The hydrogen absorption, desorption and disproportionation behaviour of the SmFe and SmFeTa alloys was observed using Thermopiesic Analysis (TPA) and Gas Flow Analysis (GFA). The difference between these two methods is in the measured quantity. In the case of TPA it is the pressure of hydrogen, and with the GFA it is the differential hydrogen flow. Schematic diagrams of both pieces of equipment are shown in **Figure 1**. The TPA is a device, which measures the changes in pressure with the sample held within a fixed volume. As the material absorbs H₂ the pressure is observed to fall, subsequent hydrogen desorption causes a pressure increase. It is important to note that these changes take place against a constantly increasing background of pressure due to the increasing temperature of the experiment. The GFA, in contrast, is a constant pressure system. Gas is allowed to flow into the system at a constant rate, typically 50 ccm/min, this flow is set by a mass flow controller. The exhaust gas flow is measured in a similar way and by taking the difference between the flow-in and the flow-out it is possible to calculate ΔQ , the amount of the gas being absorbed or desorbed at any particular stage of the experiment. In both cases, experiments were undertaken between room temperature and 800°C.

In order to assess the effect of Ta on the magnetic properties of a $Sm_2Fe_{17}N_x$ type permanent magnet, materials were processed using a HDDR based procedure. The ingot material was crushed to a particle size of less than 1 mm and then reduced further by milling. The milling was carried out in an attritor mill for times of between 1 and 60 minutes, under hexane, in the inert atmosphere of a glove box. The particle size of the attritor milled powder was determined using a Fisher sub sieve sizer. An investigation of the size distribution of the powders was undertaken with a scanning electron microscope. The HDDR processing was carried out in a rotating furnace capable of operating between 1 bar over pressure and a vacuum of 10^2 mbar. The initial stages of

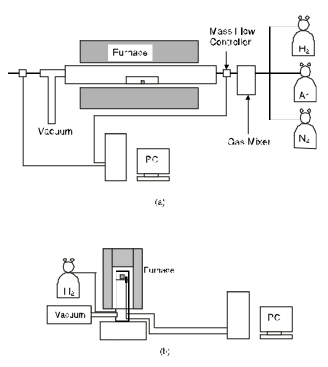


Figure 1: The schemes of GFA (a) and TPA (b) pro cesses Slika 1: Shemati~ni prikaz GFA (a) in TPA (b) procesa

the HDDR processing were carried out in pure H_2 . Samples were heated at 5°C/min to 750°C and then held at this temperature for 60 mins. The second, recombination, stage was carried out under vacuum for a further 60 mins using temperatures 740 and 820°C. All samples were subsequently nitrided at 450°C for 4 hours in a flow of nitrogen gas. Permanent magnet bonded samples were produced by mixing the powder with epoxy resin. These samples were measured at room temperature in a conventional permeameter after pulsing the magnets in a field of 5 T.

3 RE SULTS AND DIS CUS SION

3.1 X-ray diffraction experiments

Results of the lattice parameter refinement calculations can be seen in **Table 1**. These refinements reveal that the presence of Ta in the Sm_2Fe_{17} phase, in agreement with Saje et al ¹⁵, causes a significant lattice expansion. The volume of the 2:17 unit cell for the SmFeTa alloy, being in this case, some 0.42% greater than that of the binary SmFe; although this is somewhat smaller than the 0.7% expansion reported by Gutfleisch et al ¹⁶ for a 4%Nb alloy.

The changes in hydrogen pressure during heating at 5K/min, for both alloys, can be seen in **Figure 2**. The first absorption events for the binary and Ta substituted alloys are observed at temperatures between 50 and 250°C and are the result of interstitial absorption of hydrogen into the Sm_2Fe_{IT} , $SmFe_2$ and $SmFe_3$ phases. $SmFe_2$ absorbs hydrogen at temperatures between

SmFe alloy				SmFeTa alloy			
Phase	a(Å)	c(Å)	V(Å ³)	Phase	a(Å)	c(Å)	$V(Å^3)$
2:17	8.545 ± 0.992	12.428 ± 0.004	785.812±0.364	2:17	8.558±0.002	12.441±0.005	789.096±0.005

Table1: Lat tice pa ram e ter mea sure ments of 2:17 phase in SmFe and SmFeTa

 Tabela 1: Parametri osnovne celice 2:17 faze v SmFe in v SmFeTa zlitini

120-160°C and the hydride decomposes immediately to SmH_x and αFe^{17} . Because the onset temperatures for initial hydrogen absorption depend very much on the surface conditions, it would be wrong to suggest that the different behaviour of the materials is a result of the Ta substitution. Above 150°C we observe a multistage desorption. With the help of the GFA data we can interpret this multistage desorption in terms of a continuous loss of hydrogen from the 2:17 phase combined with the desorption of hydrogen from the SmFe₃H_x phase at ~245°C. The desorption from the SmFe₃H_x phase takes place over approximately 10°C after which the hydrogen loss from the 2:17 hydride continues for both materials up to ~440°C. At this point a second absorption is observed to take place until the temperature reaches ~500°C. This second absorption is clearly defined and is a consequence of the decomposition of SmFe₃H_X to SmH_X and α Fe. Further heating brings about a third desorption stage, but here there is a significant difference in the behaviour of the

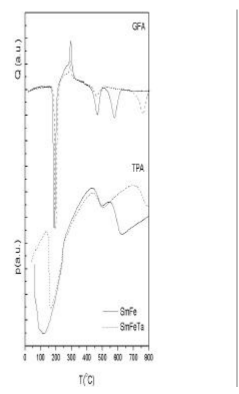


Figure 2: TPA and GFA curves for as cast SmFe and SmFeTa al loys, show ing their be hav iour during HDDR process

Slika 2: TPA in GFA krivulje za zlitini SmFe in SmFeTa, ki ponazarjata interakcije z vodikom med procesom HDDR

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two alloys. The binary material begins absorbing at ~550°C because of the $Sm_2Fe_{17}H_X$ decomposing to SmH_X and Fe whereas the Ta substituted material remains undecomposed until a temperature of ~700°C is reached.

3.3 GFA investigations

The hydrogen absorption and desorption events were also observed by measuring the difference in hydrogen flow (ΔQ) at different temperatures (Figure 2) using the GFA. The first absorption peak was observed to begin at ~170°C for both alloys. The reaction proceeds rapidly and appears to come to completion prior to the onset of the desorption stage at ~220°C. This desorption corresponds to loss of interstitial hydrogen from the 2:17 phase and at ~280°C an increased rate of desorption indicates loss of hydrogen from the SmFe₃H_X phase as well. At ~400°C we observe the beginnings of the SmFe₃ decomposition stage which takes place over the same temperature range for both materials, the process being largely completed at ~510°C. As with the TPA experiments we observe striking differences in the behaviour of the materials at higher temperatures. For the binary material the decomposition of the 2:17 phase extends from ~520°C to ~630°C, while with the Ta substituted sample it takes place from 680°C to 800°C, which is more than 150°C higher.

If we compare the results obtained with the TPA and the GFA it is clear that the different experimental methods give very comparable results. The discrepancies in the initial absorption are frequently observed and can be attributed to the variation in surface condition of the starting materials. The desorption of hydrogen from the SmFe₃H_X phase is very clearly demonstrated by the GFA and the larger peak obtained for the binary material corresponds with the relatively larger amounts of SmFe₃-type phase observed in the binary material. Both methods clearly show the decomposition of the SmFe₃H_X-type phases in both alloys between ~420 and ~520°C and the substantial difference in the temperature of decomposition for the Sm₂Fe₁₇H_X and Sm₂(FeTa) $_{17}$ H_X phases at higher temperatures.

3.4 Magnet processing

The results of a Fisher Subsieve Sizer (FSSS) particle size analysis on the SmFe and SmFeTa materials milled for between 1 and 60 mins can be seen in **Figure 3**. The measurements indicate that the average size of the

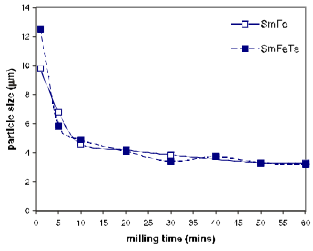


Figure 3: The part i cle size in de pend ence of mill ing time for SmFe and SmFeTa al loys

Slika 3: Velikost delcev v odvisnosti od ~asa mletja za zlitini SmFe in SmFeTa

particles decreases rapidly with milling time at least for the first 10-15 mins before settling at a size of approximately 3-4 μ m. Obviously the FSSS measurements only give us an indication of the average particle size of the powder. Information relating to the spread in the particle size is not provided by such a measurement. However, SEM micrographs of both SmFe and SmFeTa powders (Figure 4) show that the size distribution for the two materials is relatively narrow and basically similar for both materials for the same milling times.

Figure 5 shows the effect of pre-milling time on SmFe and SmFeTa processed at recombination temperatures of 740°C and 820°C. Both materials show a sharp increase in coercivity with relatively short milling times when a recombination temperature of

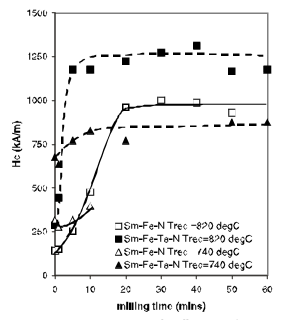


Figure 5: Variation in coercivity with milling time for SmFe and SmFeTa materials re combined at 740°C and 820°C **Slika 5:** Koercitivna sila v odvisnosti od ~asa mletja za vzorca SmFe

in SmFeTa, ki sta bila rekombinirana pri 740°C in 820°C

820°C is used. This is particularly so for the SmFeTa material. In comparison, milling time appears to have relatively little effect when the lower recombination temperature is used. In other words, while the coercivity of the materials processed at the higher temperature depends very much on the particle size of the starting material, the coercivity of the low recombination temperature samples is to a much greater extent, particle size independent. The explanation for this observation lies in the grain size of the recombined materials. The grains in the samples processed at the lower temperature

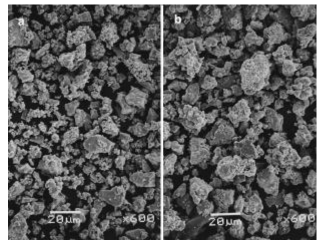


Figure 4: SEM mi cro graphs of the SmFeTa (a) and SmFe (b) pow ders milled for 20 mins

Slika 4: SEM posnetek vzorcev SmFeTa (a) in SmFe (b) mletih v attritor mlinu 20 min

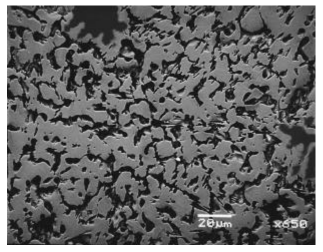


Figure 6: SEM mi cro graph show ing big grains (>20 μm) of Sm₂Fe ₁₇ phase (grey) **Slika 6:** SEM posnetek, ki prikazuje zrna Sm₂Fe ₁₇ faze (sivo)

grow relatively slowly and so the size of the particles, which contain these grains, is relatively unimportant. In the case of the higher temperature processing, where very significant grain growth can take place, particle size is critical, since grains can grow to quite large sizes in large particles, but are obviously restricted just to the size of the particle with small particles. In other words, it is impossible, for example, to have a grain size larger than 1µm in a particle with size 1µm. The 820°C recombination temperature causes rapid grain growth and the resulting coercivities in the non pre-milled (1mm) material are low as a result of this grain growth. But with a particle size reduction to 4-5µm a considerable increase in the coercivity is observed because the particle size is now in a position to restrict the grain size.

The SEM micrograph in **Figure 6** shows the presence of grains substantially larger than 5μ m in the non pre-milled starting material recombined at 820°C. Such grains could not exist in the pre-milled material simply because of the physical dimensions of the particles. This being the case, it would seem that in order to achieve the highest coercivities for SmFe based HDDR materials it is critical to restrict the grain growth by restricting the particle size.

4CONCLUSIONS

The introduction of Ta has a very significant and beneficial impact on both the cast material and the final nitrided HDDR product. The enhanced values of coercivity in the SmFeTa material indicate the critical importance of Ta in reducing as far as possible any free iron in the material. The Ta containing alloy was observed to be significantly more stable in terms of the 2:17 hydride resistance of the phase to disproportionation. This can be attributed to the dissolution of small amounts of Ta into the 2:17 phase increasing its stability with respect to decomposition in a hydrogen atmosphere.

Milling of the material before the HDDR treatment reduces the particle size and prevents the grains from

growing to a size when they may have a detrimental effect on the coercivity. The low recombination temperature samples are largely particle size independent, but the case of the high temperature processing, particle size is critical.

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