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Studies on a Novel Process for Nitrogen Utilization using Iron-based Intermetallic Compounds

(鉄系金属間化合物を用いた新規窒素利用プロセスに関する研究)

2001

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Preface

The work presented in this thesis has been carried out during 1998-2000 under the guidance of Professor Gin-ya Adachi at Department of Applied Chemistry, Faculty of Engineering, Osaka University.

This thesis deals with nitrogen absorption and desorption characteristics for iron-based intermetallic compounds and a novel reaction process using highly reactive atomic nitrogen absorbed in the intermetallic compounds. The results and findings obtained through this work gave important information on the storage technique of nitrogen and high efficient reaction using atomic nitrogen. This fundamental work is expected to contribute to the further development of the nitrogen storage materials and their applications.

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General Introduction

Metal hydrides are mainly classified to three types on the basis of the bond characters between metal and hydrogen. Intermetallic compounds such as LaNi₅ reversibly absorb and desorb large amounts of hydrogen under mild conditions around ambient temperature and pressure through the formation of interstitial type (metallic bonding type) metal hydrides [1]. For ionic and covalent bonding-types of metal hydrides such as LiH and (BeH₂)_x, the reversible hydrogen absorption and desorption can not be proceeded under the above mild conditions, since the physicochemistry is considerably different from that of the interstitial type metal hydrides. The reversible absorption and desorption of hydrogen observed on the interstitial type metal hydrides allows us the development of the hydrogen storage technique, cathode material in the Ni-H secondary battery, and so on. In addition to the reversible absorption and desorption behavior, interstitial hydrogen accommodated in the metal hydrides exists as an atomic state and it is expected to be very reactive, so that some of interstitial type metal hydrides have been found to be good hydrogenation reagents and/or catalysts [2-11].

In analogy with the metal-hydrogen system, metal nitrides are grouped into ionic bonding, covalent bonding, and metallic bonding types by the bond characters between metal and nitrogen. Among them, the rare earth-iron intermetallic compounds R₂Fe₁₇ provide the interstitial type metal nitrides by heating in N₂ or a mixed gas of NH₃-H₂ [12,13]. In particular, Sm₂Fe₁₇N₃ has been noted as one of promising materials for high-performance permanent magnets as bonded ones [14-21]. The nitrogenation for Sm₂Fe₁₇ is usually made in N₂, but this solid-gas reaction requires the much more extreme conditions of reaction pressure (~10 MPa), temperature (~550 °C) and time (~72 h) to proceed completely and uniformly than the nitrogenation in the NH₃-H₂ gas mixture [12,13]. In the latter case, large amounts of nitrogen can be interstitially incorporated in

the Sm₂Fe₁₇ crystal lattice (x=~5) under the moderate conditions (in the mixed gas of NH₃-H₂ under atmospheric pressure at ~450 °C for ~3 h) [12,13]. As a result, the excess nitrogen (Δx) in Sm₂Fe₁₇N_x (Δx =x-3) must be reversibly desorbed as N₂ down to x=3 during the annealing process by heating at ~500 °C in inert gas such as Ar to obtain the optimal nitrogen concentration for magnetic properties [12,13].

The ionic bonding type metal nitrides such as NaN₃ are commercially used as nitrogen gas generator to operate the safety bags of automobiles. Nitrogen, however, cannot be reversibly recharged due that the gas generation proceeds via the oxidative decomposition of NaN₃ (see below scheme).

$$NaN_3 \xrightarrow{KCIO_4} Na_2O + 3/2N_2$$

On the contrary to NaN₃, the nitrogen absorption and desorption behavior observed on the Sm₂Fe₁₇N_x indicates that the interstitial metal nitrides can absorb and desorb nitrogen through the formation of interstitial type metal nitrides in a similar manner as observed in the same type of metal hydrides. From this fact, it is expected that the interstitial metal nitrides are candidates for a novel nitrogen storage materials in a similar manner as the hydrogen storage ones. Furthermore, the nitrogen in the interstitial type metal nitrides exists in an atomic state and it is expected to be highly reactive. One can see that efficient nitrogenation process using highly reactive nitrogen atoms in these metal nitrides must be designed. The objects of this work are to develop a novel nitrogen storage technique using interstitial type metal nitrides and to construct new process for the nitrogenation reaction using the atomic nitrogen in these metal nitrides.

The thesis of this work consists of the following chapters:

In chapter 1, the nitrogenation and hydrogenation was conducted for the rare earth-iron intermetallic compounds R₂Fe₁₇ (R=Y, Ce, Sm) to investigate the nitrogen absorption and desorption properties over them by heating in a NH₃-H₂ mixed gas and in H₂ one, respectively. The surface of Sm₂Fe₁₇ was modified by the loading with Ru

metal, which is active for the dissociation of ammonia and molecular hydrogen. In addition, the nitrogen storage capacity was evaluated by characterizing the structure and reaction properties for these metal nitrides.

In chapter 2, the nitrogen absorption and desorption characteristics were investigated over the iron-based transition metal intermetallic compounds, viz. Laves phase-type MFe₂ (M=Ti, Zr, Nb, Mo), CsCl-type TiFe, and σ -phase VFe by heating them in the NH₃-H₂ mixed gas for nitrogenation and H₂ gas for denitrogenation. Moreover, a novel process for the nitrogenation using highly reactive nitrogen atom in the MFe₂N_x was proposed by using amorphized TiFe₂N_x powders loaded with a Ru-Al₂O₃ catalyst in N₂.

In chapter 3, the nitrogen absorption and desorption properties were investigated for the rare earth-iron intermetallic compounds, RFe₂ and RFe₇, besides on the R₂Fe₁₇-type compounds. In particular, RFe₇-type compounds belong to the same type of crystal structure for the R₂Fe₁₇ (R=Ce to Tb), where the Fe(6c) sites are partially deficient instead of the substitution of rare earth metal in 3a site, meanwhile Fe(6c) sites are fully occupied for the R₂Fe₁₇-type compounds. The differences on the nitrogen absorption and desorption properties between R₂Fe₁₇ and RFe₇ were discussed on the basis of their structural properties.

List of Publications

 Formation and Hydrogenation of the Nitrogen Stored in Iron-based Intermetallic Compounds

Ken-ichi Machida, Masahiro Itoh, Kazuhiro Hirose, and Gin-ya Adachi, Chemistry Letters, 1117 – 1118 (1998).

2) Nitrogen Storage Properties based on Nitrogenation and Hydrogenation of Rare Earth-Iron Intermetallic Compounds R₂Fe₁₇ (R=Y, Ce, Sm)

Masahiro Itoh, Ken-ichi Machida, Hiroharu Nakajima, Kazuhiro Hirose, and Gin-ya Adachi,

Journal of Alloys and Compounds, 288, 141-146 (1999).

3) Nitrogenation and Hydrogenation Characteristics of Transition Metal-Iron
Intermetallic Compounds

Masahiro Itoh, Ken-ichi Machida, Kazuhiro Hirose, Takao Sakata, Hirotaro Mori and Gin-ya Adachi,

The Journal of Physical Chemistry B, 103, 9498 – 9504 (1999).

4) Nitrogen Absorption and Desorption Characteristics for CeFe₇ and Ce₂Fe₁₇
Masahiro Itoh, Ken-ichi Machida, Kazuhiro Hirose, and Gin-ya Adachi, in preparation.

Chapter 1

Nitrogen Storage Property through Nitrogen Absorption and Desorption for Rare Earth-Iron Intermetallic Compounds R₂Fe₁₇ (R=Y, Ce, Sm)

1.1 Introduction

Rare earth-iron intermetallic compounds, R₂Fe₁₇, provide the corresponding interstitial type metal nitrides (R₂Fe₁₇N_x) by nitrogenation in N₂ or NH₃-H₂ mixed gas. Usually, the nitrogenation for R₂Fe₁₇ is made in N₂, but it needs the much more severe conditions of high pressure, temperature, and time than the nitrogenation in the NH₃-H₂ mixture gas. In the latter case, nitrogen can be easily absorbed in the R₂Fe₁₇ crystal lattice under the more moderate reaction conditions. And then, the excess nitrogen in R₂Fe₁₇N_x (x>3) is reversibly desorbed by heating in Ar atmosphere to form the R₂Fe₁₇N₃ compounds in a similar manner as metal hydride systems such as LaNi₅H_x. This behavior of nitrogen absorption and desorption has a close relation to the sites for nitrogen in the crystal lattices. Crystal structures of Sm₂Fe₁₇N_x and Y₂Fe₁₇N_x are shown in Figs. 1.1 [22]. The crystal structure of Sm₂Fe₁₇ is classified as a Th₂Zn₁₇-type one with a space group of $R\overline{3}m$ and is formed when rare earth metals (R) are Ce, Pr, Nd, Sm, Gd and Tb. Meanwhile, the R₂Fe₁₇ compounds of R = Y, Dy, Ho, Er, Tm, and Lu crystallize in another Th₂Ni₁₇-type structure (space group = P6₃/mmc) [22]. Buschow et al. [23] have concluded from neutron diffraction measurements of Th₂Fe₁₇N₃ with the same as the Th₂Zn₁₇-type structure that the nitrogen occupies the 9e sites represented by the space group $R\bar{3}m$ symmetry. However, the 9e sites can only accommodate three nitrogen atoms per an R₂Fe₁₇ formula unit and it has also deduced from the neutron diffraction measurement of Th₂Fe₁₇D_x (x= \sim 5) that the excess nitrogen in Sm₂Fe₁₇N_x

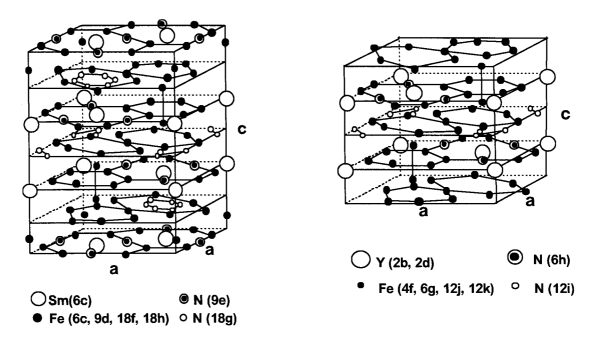


Fig. 1.1. Crystal structures of Sm₂Fe₁₇N_x (left) and Y₂Fe₁₇N_x (right).

(3<x<7) occupies other sites such as 18g site [23]. In the case of $Y_2Fe_{17}N_x$, the nitrogen shares 6h sites of the Th_2Fe_{17} -type crystal cell up to x=3, and the excess nitrogen occupies 12i sites [24]. Therefore, the nitrogen absorbed more than x=3 as $R_2Fe_{17}N_x$ is released down to x=3 since the nitrogen on the 9e or 6h sites is stable but not on the 18g or 12i sites. This behavior suggests that the R_2Fe_{17} compounds store large amounts of nitrogen in the crystal lattices in a similar manner as hydrogen storage materials. In addition, as the nitrogen in $R_2Fe_{17}N_x$ exists as a atomic state [24], and it is expected that the nitrogen in $R_2Fe_{17}N_x$ is considerably reactive, the nitrogen atoms stored above x=3 in the crystal lattice are regenerated as ammonia by replacing the atmosphere from Ar to H_2 in the above-mentioned treatment process for nitrogen desorption.

In this chapter, the nitrogenation and hydrogenation of R₂Fe₁₇ were performed in streams of NH₃-H₂ mixture and H₂ to characterize the nitrogen absorption and desorption property based on the formation of R₂Fe₁₇N_x, together with composite materials, Ru/Sm₂Fe₁₇, of which Ru metal was loaded on the surface in order to promote the reactions for nitrogenation and hydrogenation. A novel nitrogen storage

technique is proposed from the discussions on the nitrogen absorption and desorption properties over R₂Fe₁₇.

1.2 Experimental

The intermetallic compounds R₂Fe₁₇ (R= Y, Ce and Sm) were prepared from appropriate amounts of rare earth metal ingots or shots (>99.9 % in purity) and Fe rods (>99.95 % in purity) by arc melting, and then annealed in He at 1100 °C for 48 h. The ingots obtained were ground into powders with a particle size below 50 μm (BET surface area=0.15-0.30 m²/g). Further, some of Sm₂Fe₁₇ was activated by the loading of Ru metal cluster particles derived from Ru₃(CO)₁₂ on the surface of it: After 1.0 g of the Sm₂Fe₁₇ powder was impregnated with a hexane solution containing 0.06 g of Ru₃(CO)₁₂, the resulting composite materials were dried in N₂ at room temperature over night and heated at 300 °C in H₂ for 3 h.

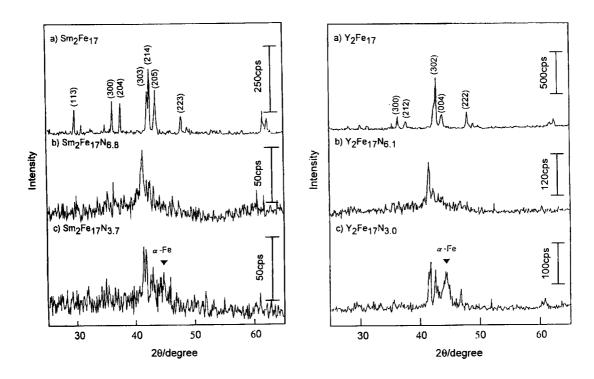
The R₂Fe₁₇ or Ru/R₂Fe₁₇ powder (1.0 g) charged in a conventional fixed-bed quartz tube reactor (12 mm o.d.) was nitrogenated in a stream of mixed NH₃-H₂ gas (molar ratio=1:1) with a flow rate of 30ml/min at 350-450 °C for 3 h (nitrogen absorption step), followed by the subsequent hydrogenation process performed in a H₂ stream with the same flow rate at 450 °C for 3 h as the absorption step. The nitrogen storage capacity of R₂Fe₁₇ powders was evaluated from the difference between the nitrogen contents directly measured on a nitrogen/oxygen analyzer (Horiba, EMGA 550) for the metal nitrides before and after the nitrogen desorption (R₂Fe₁₇N_x and R₂Fe₁₇N_{x-δ}). The structural change was checked every step for the nitrogenation and hydrogenation treatment by measurements of powder X-ray diffraction (XRD) by a MAC Science M18XHF-SHA diffractometer using Cu Kα radiation equipped with a curved graphite monochlometer. The nitrogen species (NH₃) stripped from the metal nitrides by H₂ in the nitrogen desorption step was accumulated in a cold trap (-196 °C)

and then it was analyzed on a mass spectrometer. The amount of ammonia product was quantified by the difference between the pH values of an H_2SO_4 solution trap before and after passing gas generated from the metal nitride in the nitrogen desorption step. Nitrogen absorption and desorption properties of the Sm and Fe metals used as the starting materials were also characterized on their powders (BET surface area= \sim 0.16 m² g⁻¹) by the same methods as used for R_2Fe_{17} .

The catalytic activities of ammonia decomposition over Sm₂Fe₁₇, Ru/Sm₂Fe₁₇ and Fe were measured by a gas chromatograph (Shimadzu GC-8A) in a temperature range from 200 °C to 700 °C. The inlet gas composition was 5% NH₃ and He as a balance with a flow rate of 20 ml/min. The decomposition yield from NH₃ to N₂ over each powder was calculated by integrating their peak area ratio.

1.3 Results and discussion

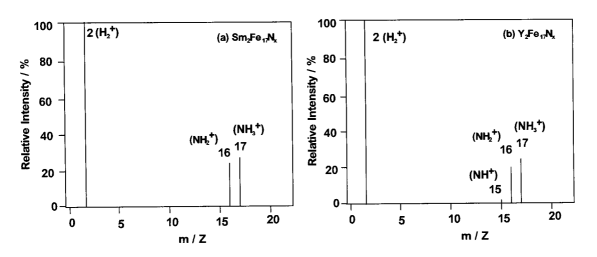
Figures 1.2 show the typical XRD patterns measured on the powdered Sm₂Fe₁₇ and Y₂Fe₁₇ nitrogenated in NH₃-H₂ mixed gas at 450 °C for 3h and hydrogenated in H₂ gas at 450 °C for 3h, together with the as-obtained one. As shown in Figs 1.2b, the diffraction peaks shifted to the lower angle side than those of the obtained after the nitrogenation treatment (nitrogen absorption). These indicate that nitrogen is incorporated interstitially into their crystal lattices as accompanied with lattice expansion. After the subsequent hydrogenation (nitrogen desorption), their XRD patterns inversely shifted to the higher angle side, suggesting lattice shrinkage for desorbing nitrogen from their crystal lattices (see Figs 1.2c). A part of the R₂Fe₁₇N_x compounds decomposed after the hydrogenation to α-Fe as shown in Figs. 1.2c because the resulting metal nitrides were metastable. The intensities of diffraction peaks for these intermetallic compounds were weaken and broaden with the nitrogenation treatments. The large amounts of the nitrogen incorporated into the crystal lattices



Figs. 1.2. XRD patterns of the Sm_2Fe_{17} (left) and Y_2Fe_{17} (right) powders: a) as obtained, b) nitrogenation in NH_3 - H_2 mixed gas at 450 °C for 3h, and c) hydrogenation in H_2 gas at 450 °C for 3h.

lowered the crystallinity.

Mass spectra for the nitrogen species regenerated from the $Sm_2Fe_{17}N_x$ and $Y_2Fe_{17}N_x$ powders heated during reactions in H_2 at 450 °C for 3 h were shown in Figs.



Figs. 1.3. Typical mass spectra for the nitrogen species eluted from (a) $Sm_2Fe_{17}N_x$ and (b) $Y_2Fe_{17}N_x$ powders with H_2 .

1.3. The mass signals were appeared at the M/Z values of 2, 15, 16 and 17, which were

assigned to H₂⁺, NH⁺, NH₂⁺ and NH₃⁺ ion species, respectively. Since NH₃ is responsible for the fraction ion species of NH⁺, NH₂⁺ and NH₃⁺, and H₂⁺ is mainly originated from the H₂ stripping gas used here, it is concluded that the nitrogen absorbed in the R₂Fe₁₇ crystal lattices is regenerated as NH₃ by stripping with H₂. In addition, the amounts of ammonia evaluated from the difference between the pH values of H₂SO₄ solution trap observed before and after the heat treatment in H₂ were in good accordance with the amount of nitrogen desorption determined directly by the nitrogen analysis for the metal nitride powders. It can be seen from these results that the nitrogen released from the R₂Fe₁₇N_x powders were converted to ammonia at almost 100% of yield.

The nitrogen absorption and desorption characteristics of R₂Fe₁₇, Ru/Sm₂Fe₁₇, Sm and Fe powders observed on the nitrogenation in the NH₃-H₂ mixed gas at 350 or 450 °C for 3 h and on the hydrogenation in H₂ at 450 °C for 3 h are summarized in Table 1.1. Even for the nitrogenation at 350 °C, Ru/Sm ₂Fe₁₇ absorbed a significant amount of nitrogen to form Ru/Sm₂Fe₁₇N_{4.9}. The nitrogen content decreased down to x=2.6 after the subsequent desorption reaction. On the other hand, no Ru metal loaded Sm₂Fe₁₇ powder was nitrogenated up to x=3.1 and desorbed nitrogen down to x=2.0 under the same reaction conditions. The nitrogen absorption-desorption property over Ru/Sm₂Fe₁₇ was superior to that over Sm₂Fe₁₇ due that the catalytic activity of Ru metal for the dissociation of NH₃ and H₂ [25]. For the nitrogen desorption step, it is considered that the hydrogen atoms diffuse into the crystal lattice and push the nitrogen in the crystal lattice out of the metal. From a calculation based on the difference of nitrogen content (Δx=2.3) between Ru/Sm₂Fe₁₇N_{4.9} and Ru/Sm₂Fe₁₇N_{2.6}, the quantity of the nitrogen desorbed from 1 cm³ of the metal nitride (p=ca.7.6 g cm⁻³) was equivalent to 13.13 mmol of NH₃, because the atomic nitrogen stored in the crystal lattice were regenerated as NH₃ at 100% of yield in the desorption step. It is noted that this amount of the nitrogen regenerated from 1 cm³ of the metal nitride is comparable to the storage

Table 1.1 Nitrogen absorption and desorption characteristics of R_2Fe_{17} , Ru/Sm_2Fe_{17} , Sm, and Fe

Compound	Cycle number	Temp./°C ^{a)}	Nitrogen content (x) b)		Regenerated
			After absorption	After desorption	ammonia (mmol/cm³) ^{c)}
Sm ₂ Fe ₁₇ N _x	1	350	3.1	2.0	6.25
$Ru/Sm_2Fe_{17}N_x$	1	350	4.9	2.6	13.13
$Y_2Fe_{17}N_x$	1	450	6.1	3.0	17.05
$Ce_2Fe_{17}N_x$	1	450	6.6	3.7	16.96
Sm ₂ Fe ₁₇ N _x	1	450	6.8	3.7	17.59
SIII2FE17IN _X	11	450	8.4	2.9	31.61
Ru/Sm₂Fe ₁₇ N _x	1	450	7.5	3.6	22.41
	11	450	8.6	2.8	36.43
SmN _x	1	350	0	0	0
	1	700	1.0	0.98	1.07
EoN	1	350	0.01	0	1.43
FeN _x	11	450	0.14	0.01	18.39

⁽a) Reaction temperature of nitrogenation.

capacity of conventional high-pressure nitrogen containers charged at 15 MPa (6.12 mmol of N₂ gas per 1 cm³ of their containers' volume). The diffusion rate of nitrogen at 350 °C in the R₂Fe₁₇ crystal lattice, however, is not enough to form the metal nitrides uniformly, and the core region of R₂Fe₁₇ particles has not yet completely nitrogenated only by heating for 3 h. Therefore, the complete absorption of nitrogen may need the much more time for the nitrogenation than 3 h.

The nitrogenation at 450 °C absorb the more amount of nitrogen in the R₂Fe₁₇ crystal lattice than at 350 °C, due to the enhancement of diffusion rate of nitrogen and of ammonia decomposition over their surfaces. Amounts of the ammonia regenerated from the R₂Fe₁₇N_x and Ru/Sm₂Fe₁₇N_x powders were evaluated to be 16.96-17.59 and 22.41 mmol per 1 cm³ of the intermetallic compounds respectively, and these values were 1.4

⁽b) Nitrogen contents were represented per chemical formula unit by determining from the nitrogen analysis data.

⁽c) The amount of ammonia regenerated is calculated as that nitrogen released from 1 cm³ of metal nitride is converted into ammonia. For example, the NH₃ amount generated from 1 cm³ of nitrogen gas container stuffed at 15 MPa is calculated to be 12.23 mmol if the N₂ gas might be converted to NH₃ at 100% of yield.

to 1.8 times larger than the nitrogen atom content as N₂ gas stored per unit volume of the conventional high-pressure nitrogen containers (12.24 mmol). It was not observed the obvious differences in the amounts of ammonia regeneration among the individual intermetallic compounds. It might be understood that the difference between the crystal Th₂Zn₁₇-type structure for Ce and Sm and the Th₂Ni₁₇-type structure for Y has little influence on the nitrogen storage capacities of them for the similarity in their crystal structures, and the standard enthalpy values of nitride formation are almost same as one another for the rare earth metals in this study [26].

On the other hand, since the Sm metal used as one of the starting materials was inactive for the dissociation of NH₃ even at 450 °C, only a trace of nitrogen was introduced into the crystal lattice. The formation of SmN required the higher reaction temperature of 700 °C. Fe metal also hardly reacted with NH₃ at 350 °C, although a large amount of nitrogen is incorporated into their lattices for R₂Fe₁₇. At 450 °C, however, a large amount of nitrogen was introduced in the Fe metal bulk as FeN_x, and a significant amount of NH₃ (18.39 mmol per 1 cm³ of FeN_x) was regenerated by the subsequent hydrogenation. The XRD pattern of nitrogenated Fe metal powder was assigned according to that reported on an interstitial Fe₄N compound [27]. These results suggest that the reversible absorption and desorption of large amounts of nitrogen over R₂Fe₁₇ is contributed to the combination of the Sm and Fe metals, where Sm plays a role for a high nitrogen storage ability, while Fe for ability of easy nitrogen desorption.

Figure 1.5 shows the decomposition properties of ammonia over Sm₂Fe₁₇, Ru/Sm₂Fe₁₇, and Fe powders. The surface areas of these powders are 0.15-0.30 m²/g measured by the conventional BET method and nearly the same. The initial temperature of ammonia decomposition over Sm₂Fe₁₇ powder is lower than that over the Fe powder. This result indicates that the activity for ammonia decomposition over the Sm₂Fe₁₇ powder is higher than that of the Fe powder. Furthermore, ammonia was decomposed on the Ru/Sm₂Fe₁₇ composite powder at a lowered temperature compared with no Ru

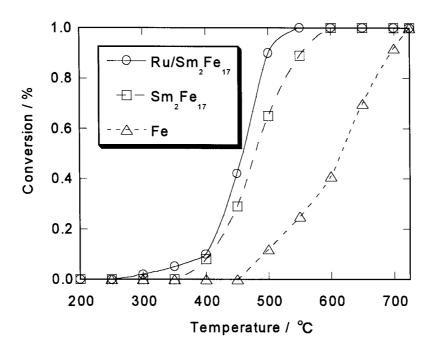


Fig. 1.4. Decomposition properties of ammonia over Sm_2Fe_{17} , Ru/Sm_2Fe_{17} and Fe powders.

metal loaded Sm₂Fe₁₇ because of the high catalytic activity of Ru metal for the NH₃ dissociation and recombination [25]. The ammonia decomposition characteristics over Sm₂Fe₁₇, Ru/Sm₂Fe₁₇ and Fe powders fairly agreed with their nitrogen storage properties summarized in Table 1. Since the catalytic activity of Fe metal and Fe₄N is lower than that of R₂Fe₁₇ and R₂Fe₁₇N_x, the Fe metal cannot absorb the large amount of nitrogen compared with them even by the nitrogenation at 450 °C.

The nitrogen storage capacity of $Sm_2Fe_{17}N_x$ was increased with repeating the nitrogen absorption-desorption cycle. An amount of the ammonia regenerated from the $Sm_2Fe_{17}N_x$ and $Ru/Sm_2Fe_{17}N_x$ powders in the 11th cycle (31.61 and 36.43 mmol) was considerably higher than that in the 1st cycle (17.59 and 22.41 mmol) (see Table 1). However, the XRD pattern of $Sm_2Fe_{17}N_x$ was mixed with those of α -Fe and amorphous phases after several nitrogen absorption-desorption cycles. This finding means that a kind of composite material, FeN_x/a -(Sm, Fe)Ny, which is derived from Sm_2Fe_{17} by repeating the nitrogen absorption-desorption cycle, gives an excellent nitrogen

absorption and desorption property, as the XRD pattern of SmN has not yet been observed on the composite sample. It is concluded that the FeN_x/a-(R, Fe)N_y composite material is also highly active for the dissociation of NH₃. In the study of ammonia synthesis using rare earth intermetallics, rare earth-iron intermetallics generated the precipitation of α -Fe after the reaction of N₂-H₂ mixed gas under a pressure of 7MPa and high catalytic activity was observed over them [28]. The increase of ammonia amount obtained in 11th cycle is ascribed to the increase of the surface area due to the crack growth on the surface and decomposition of the Sm₂Fe₁₇ by repeating nitrogen absorption-desorption cycle and the high activity of ammonia dissociation over the FeN_x/a-(R, Fe)N_y composite materials.

Schematic models for the nitrogen absorption and desorption steps took place on a $Ru/Sm_2Fe_{17}N_x$ or $Ru/Sm_2Fe_{17}N_{x-\delta}$ particle are shown in Figs. 1.5. Nitrogen is reversibly absorbed and desorbed by repeating the nitrogenation and hydrogenation cycle. A novel nitrogen storage technique can be designed by using the interstitial type metal nitrides.

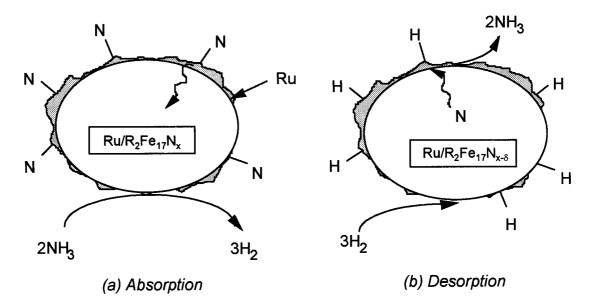


Fig. 1.5. Schematic models of the nitrogen absorption and desorption steps over the Ru/R_2Fe_{17} .

1.4 Conclusions

Rare earth-iron intermetallic compounds R₂Fe₁₇ (R=Y, Ce and Sm) absorb and desorb nitrogen reversibly through the nitrogenation by heating in the mixed gas of NH₃-H₂ and denitrogenation by heating in H₂ gas. The nitrogen storage capacities per unit volume through absorption-desorption for these samples are superior to those of conventional high pressure nitrogen containers charged at 15 MPa. More improvement for the nitrogen storage capacity of R₂Fe₁₇ than the above one is attained by loading the Ru metal on their powder surface because of the acceleration for the nitrogenation and hydrogenation. The atomic nitrogen incorporated in the compounds is regenerate as ammonia in the nitrogen desorption step at almost 100% of yield due to the high reactivity of atomic nitrogen. In addition, the nitrogen storage capacity of Sm₂Fe₁₇N_x is increased with repeating the nitrogen absorption/desorption cycle due to the formation of FeN_x/a-(Sm, Fe)N_y fine particle composites.

Chapter 2

Hydrogenation and Nitrogenation Characteristics for Iron-based Transition Metal Intermetallic Compounds

2.1 Introduction

Intermetallic compounds such as LaNi₅ have interstitial vacancies to accommodate small atoms such as hydrogen, nitrogen, and carbon. These compounds form the interstitial-type compounds by hydrogenation, nitrogenation, and carbonation. In particular, interstitial type metal hydrides can reversibly absorb and desorb large amounts of hydrogen under mild conditions. In chapter 1, it was elucidated that rare earth—iron intermetallic compounds R₂Fe₁₇, which have interstitial site vacancies, can absorb large amounts of nitrogen by heating in a flow of NH₃–H₂ mixed gas and the nitrogen incorporated into the crystal lattice is reversibly released by the subsequent heat treatment in a flow of H₂ gas. This facts indicate that rare earth-iron intermetallic compounds behave as so-called "nitrogen storage materials". Indeed, the density of the nitrogen stored in these compounds is 2-3 times higher than those for the conventional nitrogen containers charged at 15 MPa per unit volume.

On the other hand, it is known well that intermetallic compounds between transition metals such as TiFe and TiMn_{1.5} are good hydrogen storage materials [29, 30]. These compounds have enough vacant sites in crystallographic cell to accommodate hydrogen, so that it is expected that nitrogen can be also reversibly absorbed and desorbed on them like as observed on R₂Fe₁₇.

In this chapter, a series of the Laves-phase MFe₂ (M=Ti, Zr, Nb, and Mo) and AB-type M'Fe (M=Ti and V) compounds were prepared, and the nitrogen absorption and desorption properties were characterized by heating them in the NH₃-H₂ mixed gas and H₂ gas, respectively. Furthermore, the direct conversion from molecular nitrogen to

ammonia over the amorphous-like $TiFe_2N_x$ was conducted by loading the Ru/Al_2O_3 catalyst on the surface.

2.2 Experimental

The MFe₂-type compounds were prepared by arc-melting appropriate amounts of transition metal shots or ingots (>99% in purity) and Fe wire (>99.9% in purity) together in Ar (99.999% in purity). The ingots of intermetallic compounds obtained were annealed in He (99.999% in purity) at 1100 °C for 48 h and then they were crushed and pulverized into fine particles with diameters below 50 μm. The M'Fe-type compounds were prepared in the same way. A part of the TiFe₂ powder were impregnated with Ru₃(CO)₁₂ and Al(C₂H₅)₃ in their n-hexane solutions as precursors to prepare the Ru/Al₂O₃/TiFe₂ composite powder: Al₂O₃ was initially formed on the surface of the TiFe₂ particles by the oxidative decomposition of Al(C₂H₅)₃ in air at room temperature after impregnating, while Ru metal was loaded *via* the decarbonylation of the Ru₃(CO)₁₂ metal cluster compound as impregnated on the Al₂O₃/TiFe₂ composite particles by heating in H₂ at 300 °C. The procedural details were described elsewhere [31, 32].

The intermetallic compounds prepared in this study were identified on the basis of the powder X-ray diffraction measurements by a MAC Science M18XHF-SHA diffractometer using Cu Kα radiation equipped with a curved graphite monochlometer. In addition, selected area electron diffraction measurements were performed with a Hitachi H-800 transmission electron microscope operated at 200 kV. Specific surface area values for the powdered intermetallic compounds were measured by a conventional BET method, together with those of Ti and Fe powders.

Each powder of MFe₂, M'Fe, Ti, and Fe was charged in a conventional fixed-bed quartz reaction tube (12 mm o.d.), and the nitrogenation (nitrogen absorption) reactions were performed at 450-500 °C in a stream of the NH₃-H₂ mixed gas (molar ratio =1:1)

or N₂ gas with a flow rate of 30 ml/min. The hydrogenation (nitrogen desorption) reaction was carried out at 450 °C in a H₂ gas stream with the same flow rate. Their nitrogen storage properties were valuated from the differences between the nitrogen contents before and after the hydrogenation. The nitrogen content was determined by a nitrogen/oxygen analyzer (Horiba, EMGA 550). Nitrogen species eluted from the metal nitrides in the hydrogenation step were collected in a liquid nitrogen trap and analyzed on a mass spectrometer.

Catalytic activity of TiFe₂ and Fe for ammonia decomposition was estimated by monitoring the integrated peak intensity ratio between NH₃ and its degradation product (N₂) measured on a gas chromatograph (Shimadzu GC-8A) in a temperature range of 350-700 °C. The inlet gas composition was 5% NH₃ and He was used as a balance with a flow rate of 20 ml/min.

2.3 Results and discussion

2.3.1 Amorphous-like modifications induced by nitrogen absorption

From the XRD measurements, it was confirmed that all the intermetallic compounds prepared in this study were formed in a single phase except for the case of TiFe. The XRD patterns of TiFe₂, NbFe₂, and MoFe₂ were assigned to the C14 hexagonal Laves-phase structure while that of ZrFe₂ to the C15 cubic Laves-phase one. Figure 2.1 shows the XRD patterns for the respective TiFe₂ powders as nitrogenated at various temperatures (350-700 °C) in the NH₃-H₂ mixed gas for 3 h and subsequently hydrogenated at 450 °C in H₂ gas for 3 h, together with the virgin TiFe₂. As shown in Fig. 1b, the diffraction peaks for the TiFe₂ powder nitrogenated at 350 °C shifted to the lower angle side than that of the virgin specimen. This indicates that nitrogen was shared the interstitial sites to expand the crystal lattice. The lattice parameters for the TiFe₂ and TiFe₂N_{0.53} (nitrogenated at 350 °C) compounds were determined by the least squares method as follows: α =4.793 and c=7.822 Å: α =4.820 and c=7.835 Å,

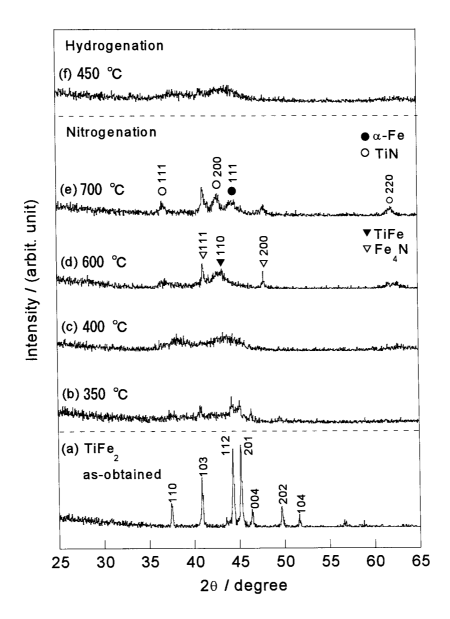


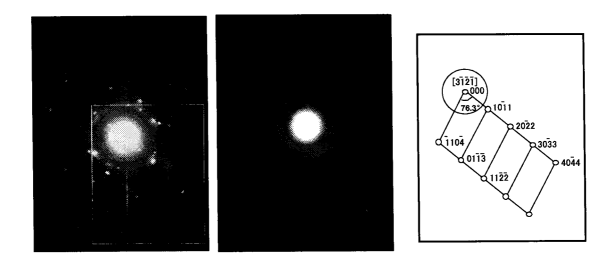
Fig. 2.1. X-ray diffraction patterns observed on the TiFe $_2$ powder nitrogenated at various temperatures (350-700 °C) in NH $_3$ -H $_2$ mixed gas for 3 h (b-e) and hydrogenated in H $_2$ gas at 450 °C for 3 h (f), together with the as-obtained one (a). The diffraction patterns for the nitrogenated and hydrogenated TiFe $_2$ are shown below and above dashed line, respectively.

respectively. On the other hand, the TiFe₂ powder as nitrogenated at 400 °C absorbed nitrogen up to TiFe₂N_{1.44} and only provided flattened broad XRD peaks. This fact suggests that the crystalline TiFe₂ transforms into the amorphous-like one caused by incorporating large amount of nitrogen into the crystal lattice. This amorphous-like

structure was maintained after the successive hydrogenation at 450 °C for nitrogen desorption step (see Fig. 2.1f). Such amorphization also takes place for the $Sm_2Fe_{17}N_{9.9}$ compound as reported by Köeninger *et al.* [33], but the resulting amorphous-like material is recrystallized to form the α -Fe phase by the hydrogenation at 450 °C. The nitrogenation at 600 °C resulted in the decomposition of the TiFe₂ compound, and additional XRD patterns assigned to Fe₄N and TiFe appeared. Further decomposition occurred during the nitrogenation at 700 °C to produce TiN, Fe₄N, and α -Fe phases. Therefore, the decomposition of TiFe₂ during nitrogenation is elucidated to take place according to the following scheme:

$$TiFe_2 \xrightarrow{in\ NH_3\ 600^{\circ}C} TiFe, Fe_4N \xrightarrow{in\ NH_3\ 700^{\circ}C} TiN, Fe_4N, Fe_4N$$

Figures 2.2a and b show selected-area electron diffraction patterns for the crystalline and amorphous TiFe₂, respectively. The net pattern shown in Fig. 2a was consistently indexed as that of [3 1 2 1] zone axis pattern of TiFe₂ with the C14 structure (see Fig. 2.2c). On the contrary, the TiFe₂ specimen nitrogenated in NH₃-H₂ at 450 °C did not show any obvious net pattern and instead a broad halo pattern composed of diffuse rings appeared which was characterized as the amorphous-like TiFe₂ phase (see Fig. 2.2b).



Figs. 2.2. Selected area electron diffraction patterns for (a) the crystalline $TiFe_2N_{0.53}$, (b) amorphous-like $TiFe_2N_{1.44}$, and (c) indexed key diagram for the diffraction pattern shown in (a).

Therefore, one can see that the electron diffraction patterns obtained are entirely consistent with the flattened broad profiles recorded on the XRD measurements.

Similar amorphization was observed in the C14 and C15 Laves-phase compounds such as ZrFe2, NbFe2, and MoFe2 during the nitrogenation at 450 °C. These flattened XRD patterns suggest that the Laves-phase compounds are easily amorphized by the nitrogenation, although no hydrogen-induced amorphization has been reported on them [34]. For the hydrogen-induced amorphization observed on the C15 Laves-phase AB₂ compounds, the atomic radius ratio between the A and B elements is the most critical factor to determine whether such amorphization takes place or not [35]. The amorphization occurs only in the C15 Laves-phase compounds with the atomic radius ratios between the A and B elements (R_A/R_B) above 1.37. An ideal R_A/R_B value is 1.225 over all the Laves-phase compounds, but most of them are not in such ideal ratio value. Therefore, the crystalline Laves-phase compounds are usually stabilized by enlarging and contracting atomic radii for the individual A and B elements. The R_A/R_B value above 1.225 results in a lower stability of the Laves-phase structure, and the strained crystalline phase are consequently forced to collapse and transformed into such amorphous phase by the insertion of hydrogen into their lattices. The crystal structures for the Laves-phase compounds consist of quite analogous layered networks. They are classified into C14 MgZn₂-, C15 MgCu₂-, and C36 MgNi₂-types by the stacking manner of layered networks. The lattice hydrogen absorbed in the C14 Laves-phase compounds occupies the tetrahedral sites surrounded by A2B2 (two A metals and two B metals), A1B3, and B4 as well as those in the C15 Laves-phase ones [36]. It is assumed that the amorphization induced by nitrogenation can be discussed in term of the R_A/R_B value in a manner similar to the hydrogen-induced amorphization. The respective R_A/R_B values for the TiFe₂, ZrFe₂, NbFe₂, and MoFe₂ are calculated to be 1.12, 1.24, 1.14, and 1.10 by using Goldschdmit's atomic radius, and the degrees of deviation from the ideal value (1.225) for these compounds ($|\Delta R_A/R_B|$ =0.015-0.125) are smaller than those observed in the hydrogen-induced amorphization. Although the R_A/R_B value for $ZrFe_2$ is 1.24, which is responsible for its quasi-ideal Laves-phase structure and is never amorphized by hydrogenation, the nitrogenation, surprisingly, induces the amorphization. This fact indicates that the insertion of nitrogen atoms into the crystal lattice more readily induces the amorphization than that of hydrogen owing to the larger size of nitrogen atom than that of hydrogen one.

Figure 2.3a shows the XRD pattern for the virgin TiFe. It was assigned to the CsCl-type structure and an unknown phase. The crystalline TiFe did not be transformed into such amorphous-like one even after the nitrogenation (see Fig. 2.3b). The nitrogen

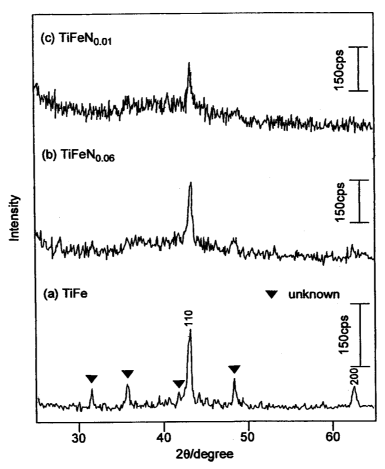


Fig. 2.3. X-ray diffraction patterns for the TiFe powders: (a), as obtained; (b), nitrogenated in NH₃-H₂ mixed gas at 450 °C for 3 h; and (c), hydrogenated in H₂ gas at 450 °C for 3 h.

content for TiFe after the nitrogenation was negligible because of the low nitrogenation rate of TiFe. This tendency is similar to hydrogenation of it [29]. In the case of VFe, however, a large amount of nitrogen was incorporated into the crystal lattice by the nitrogenation. The XRD peaks as assigned to the CrFe-type structure with tetragonal symmetry were flattened after the nitrogenation. It is presumed that the nitrogen atoms are incorporated into the tetrahedral sites surrounded by randomly or orderly occupied V and Fe atoms in the crystal lattice of CrFe-type structure [37]. These structural modifications observed on the MFe₂ and VFe indicate that the interstitial site in their crystal lattices for nitrogen occupation are too small to retain the crystalline phase, so that the nitrogen-induced amorphization more widely occurs over the intermetallic compounds compared with the hydrogen-induced one.

2.3.2 Nitrogen absorption-desorption property

Figure 2.4 shows time dependence of the heat treatments on the nitrogen content

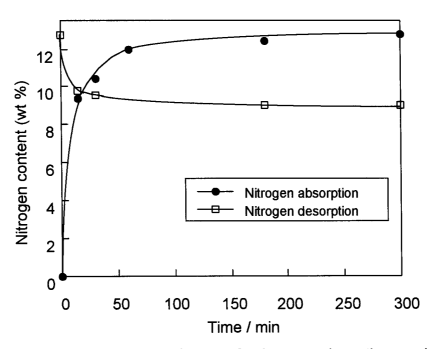


Fig. 2.4. Time dependence of nitrogen absorption and desorption observed on the TiFe₂ powder. (a) nitrogenation in NH_3 - H_2 mixed gas at 450 °C, (b) denitrogenation in H_2 gas at 450 °C.

of TiFe₂ for the nitrogen absorption and desorption processes in NH₃-H₂ and H₂ at 450 °C, respectively. The nitrogen contents of TiFe₂ in the absorption and desorption reaction steps were almost saturated at the heating time for 60 min. These results suggest that the nitrogenation and hydrogenation reaction at 450 °C for 3h are sufficient to achieve such equilibrium nitrogen contents over the TiFe₂ powder with particle diameter of 50 μm. By postulating the nitrogen absorption obeys a parabolic time law, Uchida *et al.* proposed an equation for the nitrogen diffusion to the bulk represented as [38, 39],

$$x = R\{t \exp(-A/kT)\}^{1/2}$$
 (2.1)

where x is the nitrogen concentration, A the activation energy for diffusion, k the Boltzmann constant, T the absolute temperature, and R the temperature-independent constant. Since the nitrogenation and denitrogenation reactions obey the Eq. 2.1 below the reaction time of 30 min, the activation energy of the nitrogen diffusion was calculated from an Arrhenius plot for the temperature dependence of slope $d(x^2)/dt \, vs$. the reciprocal of heating temperature to be 116.3 kJ/mol in the temperature range

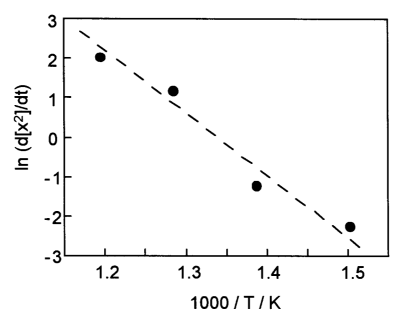


Fig. 2.5. Arrhenius plot of the nitrogen absorption rate of $TiFe_2$ in NH_3 - H_2 .

between 350 and 500 °C (see Fig. 2.5).

Figure 2.6 shows the typical mass spectrum observed on gaseous species eluted from $TiFe_2N_x$ during the hydrogenation. The mass signals at the m/z values of 2, 15, 16, and 17 were assigned to H_2^+ , NH_2^+ , NH_2^+ , and NH_3^+ ion species respectively. The H_2^+ ion is derived from the elution gas of H_2 used here. Amounts of the ammonia products were evaluated from the differences between the pH values of H_2SO_4 solution trap before and after hydrogenation, and the resulting values were in good accordance with the ammonia amounts evaluated from the decreases in the nitrogen contents of the

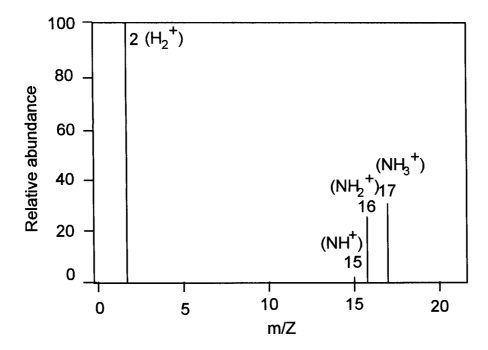


Fig. 2.6. Typical mass spectrum for the nitrogen species generated from the $TiFe_2N_x$ in the nitrogen desorption step.

hydrogenated sample directly determined by the nitrogen analysis. This result indicates that most of the nitrogen desorbed from the metal nitride is converted to ammonia in the desorption step owing to the high reactivity of atomic nitrogen.

The nitrogen absorption-desorption properties of MFe₂, M'Fe, Ti, and Fe were summarized in Table 2.1. Nitrogenation and hydrogenation treatments were performed by heating in NH₃-H₂ and in H₂ at 450 °C for 3 h, respectively. The specific surface area

Table 2.1. Nitrogen absorption and desorption characteristics of MFe₂, M'Fe, Ti, and Fe nitrogenated in the NH_3 - H_2 mixed gas

metal	cycle	nitrogen o	regenerated ammonia ^{a)}	
nitride	number	nitrogenated	hydrogenated	(mmol/cm ³ M.N.)
	1	1.48	0.93	26.25
TiFe ₂ N _x	11	1.50	0.94	26.60
ZrFe ₂ N _x	1	1.74	1.28	19.64
NbFe ₂ N _x	1	1.76	1.16	25.17
NbFe ₂ N _x	1	0.70	0.34	15.98
VFeN _x	1	0.85	0.63	15.00
TiFeN _x	1	0.06	0.01	3.10
TiN _x	1	0	0	0
FeN _x	1	0.14	0.01	18.3

a) The amount of ammonia regenerated is calculated as that nitrogen released from 1 cm³ of metal nitride is converted into ammonia.

values of the powdered samples used here are nearly same as one another (0.15-0.30 m² g⁻¹). Among them, the Laves-phase compounds, TiFe₂, ZrFe₂, NbFe₂, and MoFe₂ showed good nitrogen absorption-desorption properties *via* the transformation into the amorphous-like phases as shown in Figs. 2.1 and 2.2. The TiFe₂ compound interstitially absorbed large amounts of nitrogen into the disordered lattice up to TiFe₂N_{1.48} and, by the subsequent hydrogenation, the interstitial nitrogen was desorbed down to TiFe₂N_{0.93}. From the difference of nitrogen content (Δx =0.55) between TiFe₂N_{1.48} and TiFe₂N_{0.93}, the quantity for the nitrogen desorption per 1 cm³ of the metal nitride (ρ =*ca*. 6.86 g cm⁻³) was calculated to 26.25 mmol of NH₃, because the atomic nitrogen stored in the host lattices was regenerated as NH₃ at about 100% yield in the hydrogenation process. It is noted that the storage capacity of TiFe₂ greatly exceeds that of the conventional nitrogen container charged at 15MPa per unit volume. The amount of the ammonia

product from $1.0~\rm cm^3$ of the nitrogen gas container is equivalent to $12.2~\rm mmol$ of NH₃ if the N₂ might be converted to NH₃ in 100% of yield. The NbFe₂ and ZrFe₂ compounds also absorbed large amounts of nitrogen, which were comparable to that for the TiFe₂N_x one. Meanwhile, the amount of the nitrogen absorption for MoFe₂ was smaller and the resulting quantity of ammonia (15.98 mmol) was somewhat less than those for other Laves-phase compounds. Although the nitrogen absorption property is influenced by the factors such as the crystal structure, the nitrogen diffusion in the crystal lattices, and the catalytic activity of the surface metals, the standard formation enthalpy of metal nitride should be considered as the additional factor in this case: The enthalpy value of MoN (-92 kJ mol⁻¹) is very small compared with those for TiN (-338.1 kJ mol⁻¹), ZrN (-370 kJ mol⁻¹), and NbN (-235.1 kJ mol⁻¹).

In addition, the MFe₂ compounds showed good stability for the nitrogen absorption-desorption cycle. After the 11th nitrogen absorption-desorption cycle, the metal nitride system of TiFe₂N_{1.50} (as nitrogenated) and TiFe₂N_{0.94} (as hydrogenated) regenerated 26.60 mmol of NH₃ per 1cm³ of the metal nitride. This amount is almost same as that of the 1st cycle (26.25mmol). The XRD measurements revealed that the structure of TiFe₂N_x was still amorphous-like after 11th cycle. Similar behaviors were observed in the other MFe₂ compounds. Therefore, the Laves-phase MFe₂ intermetallic compounds can incorporate a large amount of nitrogen in their amorphous-like host lattices.

The CsCl-type compound TiFe hardly absorbed nitrogen, so that the resulting amount of ammonia product was only 3.1 mmol per 1cm³ of the metal nitride, although this compound can absorb a large amount of hydrogen. The TiFe compound has been investigated also as a catalyst for ammonia synthesis. Schwab *et al.* [40] have reported that the special pretreatment made by cycling the hydrogenation and dehydrogenation for TiFe is necessary for the formation of TiFeN_x on nitrogenation in NH₃. Indeed, Kirch *et al.* [41] have reported that the Fe/TiN composite materials of which the Fe

metal are highly dispersed in the TiN matrix via the decomposition of TiFe by nitrogenation show high activity for ammonia dissociation. On the other hand, VFe reversibly absorbed and desorbed a large amount of nitrogen, and the amount of ammonia product was 15.0 mmol per 1cm³ of the metal nitride. For the TiFe compound, it is considered that the nitrogen atoms incorporated share the octahedral sites formed by two Fe metals and four Ti metals in a manner similar to hydrogen atoms [42]. Meanwhile, the structure of VFe is classified as the σ -CrFe-type structure which is similar to β -U-type one [43]. The crystal structure of β -U consists of layered networks [44], and the isostructural analogous compounds such as VFe have many interstitial vacancies to incorporate nitrogen. It is expected that the nitrogen absorbed is located on such interstitial sites.

A large amount of ammonia product (18.3 mmol) was observed on Fe through the nitrogenation and hydrogenation. This high capacity for the nitrogen storage is also due to the formation of interstitial metal nitride. The nitrogen absorbing property of Ti, Nb, or Mo was very poor and no interstitial metal nitride phase was judged from the XRD measurements after nitrogenation. The XRD pattern for Fe after the nitrogenation consists of two phases assigned to Fe and Fe₄N. It is found that the nitrogen absorption rate of Fe powder (50 μ m) is slower than that for the TiFe₂ of similar particle diameter. However, the resulting Fe₄N phase was easily denitrogenated to α -Fe by the hydrogenation in H₂ gas at 450 °C.

Figure 2.7 shows the decomposition properties of ammonia over the TiFe₂ and Fe powders. A drastic change in the conversion from ammonia to molecular nitrogen was observed on TiFe₂ in a temperature range between 425 and 450 °C (0.5% at 425 °C to 12% at 450 °C). Meanwhile, Fe provided a steep increase in the higher temperature region (3% at 500 °C to 26% at 550 °C). This result indicates that the catalytic activity for the ammonia dissociation over TiFe₂ is higher than that over Fe, and is responsible for the large amount of nitrogen absorption for TiFe₂ as summarized in Table 1. Since

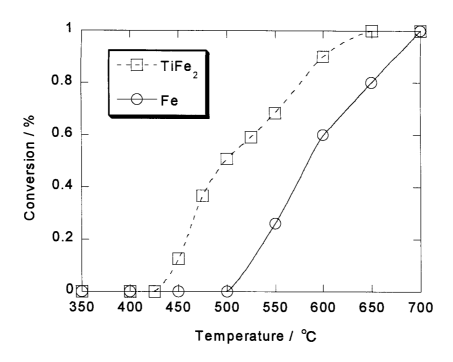


Fig. 2.7. Catalytic activity for dissociation of ammonia over the TiFe₂ and Fe powders.

the catalytic activity of Fe for the ammonia dissociation is lower than that of TiFe₂, Fe cannot be completely nitrogenated under this experimental condition, and the longer heating time is needed for the complete formation of Fe₄N.

When the molecular nitrogen is used as a nitrogen source instead of ammonia, the amount of nitrogen absorption for TiFe₂ decreased due to the high bond energy of N≡N. Aiming at promoting the catalytic activity of intermetallic compounds for the dissociation of molecular nitrogen, the Ru/Al₂O₃ catalyst as active for the dissociation and adsorption of molecular nitrogen was loaded on the surface of the TiFe₂ particles. The amounts of ammonia product for the unloaded TiFe₂ and loaded Ru/Al₂O₃/TiFe₂ powders are summarized in Table 2.2. The Ru/Al₂O₃/TiFe₂ composite powder absorbed the more amount of nitrogen compared with the unloaded one. The resulting metal nitride Ru/Al₂O₃/TiFe₂N_{0.33}, however, scarcely desorbed nitrogen in the following hydrogenation. It was found from the XRD measurements that both the TiFe₂N_{0.1} and Ru/Al₂O₃/TiFe₂N_{0.33} powders were still in the crystalline state even after nitrogenation.

Table 2.2. Nitrogen absorption and desorption characteristics for no loading and loading Ru/Al₂O₃ on the surface of crystalline and amorphous TiFe₂

metal nitride	cycle _ number	nitrogen c	regenerated			
		Nitrogenated	hydrogenated	ammonia (mmol/cm³ M.N.)		
(A) nitrogenated in the N₂ gas at 500 °C for 3 h						
c-TiFe ₂ N _x *	1	0.10	0.10	0		
a-TiFe₂N _x *	1	1.14	1.14	0		
Ru/Al ₂ O ₃ / c-TiFe ₂ N _x	1	0.33	0.33	0		
Ru/Al ₂ O ₃ / FeN _x	1	0	0	0		
	1	1.57	1.51	2.57		
Ru/Al ₂ O ₃ / a-TiFe ₂ N _x	2	1.60	1.55	2.14		
	5	1.72	1.67	2.14		
(B) nitrogenat	ed in the NH ₃ -H ₂	mixed gas at 450	°C for 3 h		
Ru/Al ₂ O ₃ / a-TiFe ₂ N _x	1	1.75	1.44	13.6		

^{*} c-TiFe2 and a-TiFe2: crystalline and amorphous TiFe2, respectively.

It is presumed that the nitrogen atom incorporated in the stable tetrahedral sites surrounded by Ti2Fe2 atom units in such low nitrogen concentration region and can not be released owing to the strong interaction between the nitrogen atom and its incorporated site. Therefore, once the Ru/Al₂O₃/TiFe₂ composite material is nitrogenated by heating in NH₃-H₂ to be amorphized, the subsequent denitrogenation allows us to regenerate ammonia by desorption of excess nitrogen stored in the amorphized lattice.

The pretreated Ru/Al₂O₃/a-TiFe₂N_x reversibly absorbed and desorbed nitrogen even though molecular nitrogen was used as the nitriding source, providing 2.57 mmol of ammonia per 1cm³ of the metal nitride. Meanwhile, a-TiFe₂ with no loading catalyst was hardly nitrogenated under the same condition. Furthermore, the Ru/Al₂O₃/Fe composite powder as prepared from the Fe powder according to the same procedure for

 $Ru/Al_2O_3/TiFe_2$ was not nitrogenated in N_2 and never produced any ammonia by the subsequent hydrogenation. These results mean that the amorphous-like phase is essential for their reversible nitrogen absorption and desorption and the Ru/Al_2O_3 catalyst plays a significant role in the dissociation of $N\equiv N$ bond.

Table 2.3. Determination of The Ammonia Product from the Ru/Al₂O₃/ a-TiFe₂N_x Composite Powder as Listed in Table 2

analysis	hydrogo	enation	generated ammonia (μmol/cm³ of M.N.) ^{a)}
	before	after	
pH value (H₂SO₄ aq.)	2.52	10.09	324

a) Amount of ammonia product is evaluated from changes of the pH value of H_2SO_4 aqueous solutions before and after introducing H_2 elution gas from the metal nitride.

The Ru/Al₂O₃/a-TiFe₂ composite material maintained good stability against the ammonia synthesis cycle. In the 5th cycle, the amount of ammonia product was evaluated from the difference between the nitrogen contents for the nitrogenated and hydrogenated powders to be 2.14 mmol per 1cm³ of metal nitrides (see Table 2.2). The quantity of nitrogen desorption is almost comparable to the 1st cycle. In addition, this value fairly agreed with the amount of ammonia product determined by monitoring the pH value of the H₂SO₄ solution trap before and after hydrogenation as listed in Table 2.3. This result implies that a novel reaction process for nitrogenation using nitrogen storable intermetallic compounds should be developed. Aika et al. [45, 46] have reported that rare earth oxide-promoted Ru metal catalysts effectively produce ammonia at atmospheric pressure with a rate of 300-800 μmol g⁻¹ h⁻¹. Meanwhile, the amount of ammonia product over the Ru/Al₂O₃/a-TiFe₂ composite material per one nitrogenation-hydrogenation cycle was 324 μmol g⁻¹. This value is not small compared with the rare earth oxide-promoted Ru metal catalyst, because the surface area of the

TiFe₂ powder used here was only 0.24 m² g⁻¹, while that of the rare earth

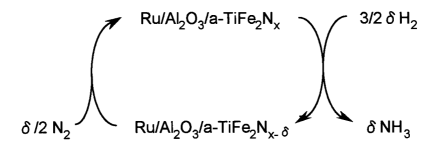


Fig. 2.8. A schematic model for a novel nitrogenation reaction using nitrogen storage material.

oxide-supported Ru metal catalyst was estimated to be 35 m² g⁻¹ from the literature at least [46]. Figure 8 shows a schematic model for new nitrogenation reaction using the composite metal nitrides as proposed in this study.

2.4 Conclusions

The crystal Laves-phase compounds MFe₂ (M=Ti, Zr, Nb, and Mo) transform to the amorphous-like phase induced by the nitrogenation in the NH₃-H₂ mixed gas at 450 °C for 3 h, although such kinds of amorphization do not occur by the hydrogenation. This fact indicates that the nitrogen-induced amorphization easily takes place compared with the hydrogen-induced one owing to the difference in the atomic radius between nitrogen and hydrogen. A similar structural modification behavior is also observed on the σ-phase VFe compound with the layered structure.

The intermetallic compounds MFe₂ and VFe reversibly absorb and desorb large amounts of nitrogen in the amorphous-like metal nitride, when ammonia is used as nitrogen source. In the nitrogen desorption step, most of the nitrogen released from the metal nitrides reacts with hydrogen to produce ammonia owing to the high reactivity of atomic nitrogen liberated from the metal nitrides. Additionally, the amorphous-like MFe_2N_x show good stability against the nitrogen absorption-desorption cycle. The $Ru/Al_2O_3/a$ - $TiFe_2N_{x-\delta}$ composite powder, which is once amorphized by the

nitrogenation in NH_3 - H_2 and subsequently hydrogenated in H_2 , can absorb nitrogen even using inert molecular nitrogen as the nitrogen source, and the resulting metal nitride releases nitrogen as ammonia by heating in H_2 at ambient pressure. The amount of ammonia product in one nitrogenation-hydrogenation cycle is about 324 μ mol g⁻¹, which allows us to develop a novel process for nitrogenation reaction using the interstitial metal nitrides.

Chapter 3

Nitrogen Absorption and Desorption Characteristics over Rare earth-Iron Intermetallic Compounds

3.1 Introduction

In chapter 1, from the nitrogenation and hydrogenation characteristics of intermetallic compounds R_2Fe_{17} (R=Y, Ce and Sm), it has been found that significant amounts of nitrogen are absorbed by heating in the NH₃-H₂ mixed gas and the resulting metal nitrides reversibly desorbed the nitrogen as NH₃ by heating them in H₂ at 450 °C. Besides R_2Fe_{17} , other rare earth-iron intermetallic compounds with chemical formula of RFe₇ and RFe₂ have been studied by some workers [47-51]. RFe₇ and RFe₂, which possess the interstitial free spaces in crystal lattices to accommodate nitrogen, are expected to absorb and desorb nitrogen in a similar manner as the R_2Fe_{17} compounds. It is noted that the atomic arrangement for RFe₇ is intermediate between those of CaCu₅- and Th₂Zn₁₇-type structures and this structure is regarded as the iron deficient Th₂Zn₁₇-type structure, so that it is expected that the nitrogen more easily diffuses in the RFe₇ crystal lattices than that in R_2Fe_{17} ones.

In this chapter, rare earth-iron intermetallic compounds were prepared. The nitrogenation and hydrogenation were performed for these compounds in NH₃ or N₂ and H₂ in a temperature range of 250-450 °C to investigate the nitrogen absorption-desorption properties. Additionally, in order to promote the dissociation of molecular nitrogen as a nitriding gas, ruthenium metal catalyst (Ru/La₂O₃) was loaded on the surface of the CeFe₇ particles. The nitrogen storage and ammonia production properties using N₂ as the nitrogenation source were studied by cycling the nitrogenation and hydrogenation processes.

3.2 Experimental

The RFe₇ (R=Ce, Nd, Sm) and CeFe₂ compounds were prepared from appropriate amounts of rare earth metal shots or ingots (>99% in purity) and Fe wire (>99.9% in purity) by the conventional arc melting method in Ar atmosphere. The ingots of intermetallic compounds obtained were annealed in He at 800 °C for 48 h, and then crushed and pulverized into fine powders with diameters below 50 μm. For a part of the CeFe₇ powder, the Ru-La₂O₃ catalyst was loaded on the surface of CeFe₇ by the mechanical grinding method. La₂O₃ was heated in a flow of H₂ gas at 600 °C for 5 h to remove the moisture before use. La₂O₃ and CeFe₇ (molar ratio, La₂O₃:CeFe₇ = 1:5) were mixed in the N₂ gas atmosphere using a planetary-type ball mill. Ru was subsequently loaded on the surface of the La₂O₃-CeFe₇ composite powder by impregnation in a Ru₃(CO)₁₂-hexane solution via the decarbonylation by heating in H₂ at 573 K.

The intermetallic compounds prepared were identified on basis of powder X-ray diffraction patterns measured on a MAC Science M18XHF-SHA diffractometer using Cu Kα radiation equipped with a curved graphite monochlometer. The surface area values for the intermetallic compound powders were measured by a conventional BET method, together with those of Fe and Ce powders. The BET surface area values of all the samples studied in this study were almost similar to one another (0.16-0.24 m² g⁻¹).

Every weighed powder (1 g) of RFe₇, R₂Fe₁₇, and RFe₂ was charged in each conventional fixed-bed quartz tube (12 mm o.d.), and the nitrogenation was made for them in the temperature range of 250-450 °C for 3 h in the respective streams of NH₃, NH₃-H₂, or N₂ gas with a flow rate of 30 ml/min. The hydrogenation was subsequently carried out for the respective nitrogenated samples at 450 °C for 3 h in the H₂ stream of the flow rate as that of NH₃ or N₂ for the nitrogenation. Nitrogen contents of nitrogenated and hydrogenated samples were measured by a nitrogen/oxygen analyzer (Horiba, EMGA 550) and the amount of the nitrogen regenerated from the metal nitride was calculated from the difference between their respective nitrogen contents. Nitrogen

species derived from the metal nitrides in the hydrogenation, that is, the nitrogen desorption step were collected in a liquid nitrogen trap and characterized on a mass spectrometer. The nitrogen regenerated as ammonia was quantified by monitoring the pH values of H₂SO₄ solution in the trap before and after passing the stripping gas from the metal nitride powder in the hydrogenation step.

The catalytic activities of ammonia decomposition over the RFe₇ and R₂Fe₁₇ powders were measured by a gas chromatograph (Shimadzu GC-8A) in a temperature range of 200 °C-750 °C. The inlet gas was 5% NH₃ and He was used as a balance gas with a flow rate of 20 ml min⁻¹. The decomposition yield (conversion) from NH₃ to N₂ over the powdered samples was calculated by integrating their peak area ratio.

3.3 Results and discussion

The XRD patterns for the RFe₇ and RFe₂ powders prepared in this study were mainly indexed on the rhombohedral symmetry with space group of $R\overline{3}m$ and C15 Laves phase with space group of Fd3m, respectively. Figure 3.1 shows the XRD patterns for CeFe₇ heated in NH₃ at 400 °C for 1 h and in H₂ at 450 °C for 3 h, together with the as-obtained one. As shown in Fig. 3.1a, the diffraction patterns of the as-obtained CeFe₇ compounds fairly agreed with the crystal data reported by Ray [47]. There are seven formula units in the crystallographic primary cell, and the crystal structure of the RFe₇ phase prepared by the arc melting consists of the intermetallic atom arrangement between the CaCu₅- and Th₂Zn₁₇-type ones. This type of crystal structure is composed of one AB₅ layer alternating with two A₂B₁₇ layers, *i. e.* 3AB₅ + 2A₂B₁₇ = 7AB₇ [47]. The XRD patterns for the RFe₇ prepared by arc melting are obviously distinct from the patterns reported elsewhere [48-50]. It seems that the structural arrangement of the resulting RFe₇ is greatly affected by the preparation methods, by the arc melting, self-flux crystal growth, rapid quenching and mechanical alloying. For the CeFe₇ sample prepared in this work, in

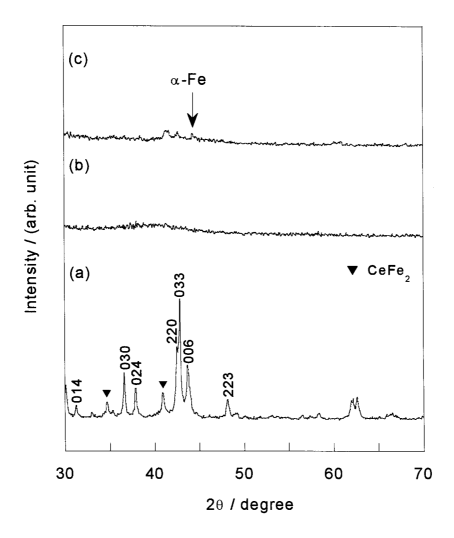


Fig. 3.1. XRD patterns of the CeFe $_7$ powders: a) untreated (as-obtained), b) nitrogenated in NH $_3$ at 400 °C for 1 h, and c) hydrogenated in H $_2$ at 450 °C for 3 h. Nitrogen contents x in the b) and c) samples are calculated to be 4.31 and 2.10 as the converted values per chemical formula unit of the starting material CeFe $_7$.

addition, weak peaks of CeFe₂ were also observed on the XRD patterns (see Fig. 3.1a). The phase diagram in the Ce-Fe binary system suggests that the CeFe₂ and CeFe₇ compounds are usually formed peritectically by the conventional arc melting method [51]. Figure 3.1b shows the diffraction pattern for RFe₇ after nitrogenation in NH₃ at 400 °C (nitrogen absorption). The diffraction peaks for the CeFe₇ powder disappeared by the nitrogenation, indicating that the crystalline CeFe₇ phase transformed to the amorphous one as seen in the Laves phase compounds such as TiFe₂. Similar amorphization was

observed on the nitrogenated CeFe₂ powder. After the hydrogenation (nitrogen desorption) in H₂ at 450 °C for 3 h for CeFe₇N_x, the weak diffraction peaks of α -Fe phase and CeFe₇N_x were observed on the XRD pattern (see Fig. 3.1c). CeFe₇N_x is the metastable compound, so that a part of CeFe₇N_x decomposed and generated the precipitation of α -Fe. Amorphous CeFe₇N_x was recrystallized by the desorption of nitrogen, although the peak intensity was considerably weak, suggesting that the crystallinity of CeFe₇N_x was very low.

Figure 3.2 shows a typical mass spectrum for the nitrogen species regenerated from the CeFe₇N_x powder by heating in H₂ at 450 °C for 3 h. The mass signals were appeared at the m/z values of 2, 15, 16 and 17, which were assigned to H_2^+ , NH^+ , NH_2^+ and NH_3^+ ions, respectively. Since H_2^+ is mainly originated from the H₂ stripping gas used here, NH₃ is responsible for the fraction ion species of NH^+ , NH_2^+ and NH_3^+ . It is concluded that the nitrogen absorbed in the RFe₇ crystal lattices is regenerated as NH₃ by reacting with H₂

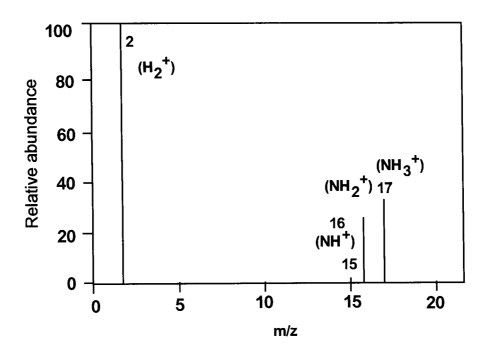


Fig. 3.2. Typical mass spectrum for the nitrogen species eluted from the amorphous (Ce, Fe)N $_{\rm x}$ with a converted composition of CeFe $_7$ N $_{3.1}$ with H $_2$.

because any signals for the ion species derived from N₂ are not observed. The amounts of ammonia product determined by monitoring the pH values of H₂SO₄ solution trap before and after passing the H₂ gas were in good accordance with the quantities of the desorbed nitrogen as directly determined for the nitrogenated and hydrogenated metal nitrides by nitrogen analysis. It can be seen from these results that the atomic nitrogen released from the nitrogenated CeFe₇ powder is converted to ammonia at about 100% of yield.

The nitrogen absorption and desorption characteristics of the CeFe₂, CeFe₇, and Ce₂Fe₁₇ powders observed on the nitrogenation in NH₃–H₂ at 450 °C for 3 h and on the hydrogenation in H₂ at 450 °C for 3 h are summarized in Table 3.1. The properties of nitrogen absorption and desorption on the Th₂Zn₁₇ type compounds were superior to the C15 laves phase type one. In Th₂Zn₁₇ type compounds, RFe₇ type compound, which is iron deficient Th₂Zn₁₇ type structure, can absorb and desorb larger amounts of nitrogen compared with R₂Fe₁₇ type compound. The smaller ammonia regeneration over the CeFe₂ powder was presumably due to the less activity of ammonia dissociation over the surface of CeFe₂, since the same type C15 Laves phase compound, ZrFe₂, can absorb a large amount of nitrogen (see Table 2.1 in Chapter 2).

The nitrogenation in NH₃ proceeds via the dissociation of ammonia over the surface

Table 3.1. Nitrogen absorption and desorption characteristics for CeFe $_2$, CeFe $_7$, and Ce $_2$ Fe $_{17}$

Metal Nitroge		content (x)	Regenerated ammonia
nitride (M.N.)	Nitrogenated	Hydrogenated	(mmol cm ⁻³ of M.N.) ^{b)}
CeFe₂N _x	1.2	0.9	11.33
CeFe ₇ N _x	3.3	0.6	28.30
Ce ₂ Fe ₁₇ N _x	6.6	3.7	16.96

a) Nitrogen contents x of the samples are represented as the values per chemical formula unit.

b) The amount of ammonia regenerated is calculated by assuming that the nitrogen released from 1 cm³ of metal nitride is converted to ammonia.

of the intermetallic compounds as following equilibrium.

$$NH_3 \Leftrightarrow 3/2H_2 + N$$
 $\Delta G^0 = 98450 - 42.3T (J)$ (3.1)

From the Eq. 3.1, the activity of atomic nitrogen is represented as following equation.

$$a_N = \frac{P_{NH_3}}{P_{H_3}^{3/2}} \exp(-\frac{\Delta G^0}{RT})$$
 (3.2)

From equation 3.2, it is obvious that the nitrogenation in sole NH₃ is more preferable than that in a NH₃-H₂ mixed gas with the consideration of the H₂ partial pressure, although H₂ in the nitrogenation by NH₃-H₂ mixed gas plays a role of easy nitrogen diffusion in the lattice due that the previous insertion of hydrogen into the crystal lattice accompanies the lattice expansion. So the nitrogen absorption and desorption characteristics of the R₂Fe₁₇ and RFe₇ powders by the nitrogenation in NH₃ at 450 °C for 3 h and hydrogenation in H₂ at 450 °C for 3 h were performed (Table 3.2). Change of nitrogenation atmosphere from NH₃-H₂ to NH₃ allows absorbing large amounts of nitrogen over both R₂Fe₁₇ and RFe₇. This result suggests that nitrogenation by sole NH₃ is more suitable than that by NH₃-H₂ mixture. The nitrogen absorption and desorption properties over the RFe₇ type

Table 3.2. Nitrogen absorption and desorption characteristics for the $R_2 \mbox{\rm Fe}_{17}$ and $R\mbox{\rm Fe}_7$

Metal	Nitrogen content (x)		Regenerated ammonia	
nitride (M.N.)	Nitrogenated	Hydrogenated	(mmol cm ⁻³ of M.N.) ^{a)}	
Ce ₂ Fe ₁₇ N _x	8.86	4.28	31.82	
$Nd_2Fe_{17}N_x$	8.06	3.56	31.25	
Sm₂Fe₁ ₇ N _x	9.01	4.43	31.82	
CeFe ₇ N _x	4.32	2.10	36.16	
NdFe ₇ N _x	4.08	1.81	33.21	
SmFe ₇ N _x	4.15	1.85	33.92	

a) The amount of ammonia regenerated is calculated by assuming that the nitrogen released from 1 cm³ of metal nitride is converted to ammonia.

compounds surpass the R₂Fe₁₇ type ones in also this nitrogenation condition. Fe (6c) site is partially occupied (occ=2/3) in the crystallographic cell for CeFe₇, whereas Fe (6c) site is fully occupied (occ=1) for Ce₂Fe₁₇. It is concluded that the difference of nitrogen absorption and desorption properties between CeFe₇ and Ce₂Fe₁₇ are contributed to the difference of crystal structure among them. If the nitrogen absorption and desorption behavior obeys a parabolic time law and reaction between metal and nitrogen is sufficient fast, the nitrogen diffusion in the bulk represented as Eq. 2.1 (see Chapter 2). It is necessary that the change of nitrogen concentration is proportional to the square root of the reaction time from equation 2.1. Figure 3.3 shows the time dependence of the amount of nitrogen desorption for CeFe₇N_{3.0} and Ce₂Fe₁₇N_{7.54} in H₂ flowing at 300 °C. As the good linearity was observed between the amount of nitrogen desorption and square root of the reaction time, the activation energies for nitrogen diffusion were evaluated from the nitrogen desorption rates for the nitrogenated specimen CeFe₇N_{3.0}

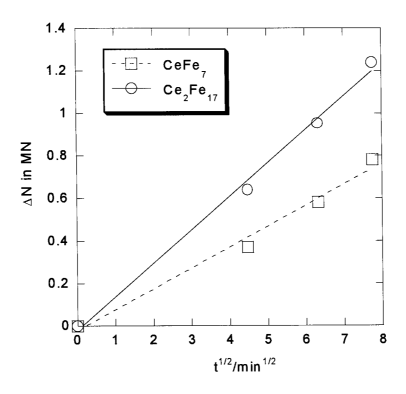


Fig. 3.3. Amount of Nitrogen desorption vs. the square root of the reaction time over $CeFe_7N_x$ and $Ce_2Fe_{17}N_x$.

and Ce₂Fe₁₇N_{7.54} in the temperature range 250-350 °C (see Fig. 3.4). The activation energies of nitrogen diffusion are 69.3 and 101.6 kJ/ mol for CeFe₇ and Ce₂Fe₁₇, respectively. The crystal structure of CeFe₇ is regarded as a kind of iron deficient Th₂Zn₁₇-type structure. Therefore, the nitrogen atoms diffuse more easily in CeFe₇ than that in Ce₂Fe₁₇ through iron-deficient sites of the former ones. One of the reasons for the superior property of nitrogen absorption and desorption observed on CeFe₇ is responsible for the ease of nitrogen diffusion in the bulk compared with Ce₂Fe₁₇.

To compare the nitrogen storage properties for NdFe₇ and Fe, the nitrogen absorption and desorption characteristics of the NdFe₇ and Fe powders by the nitrogenation in NH₃ at 300-450 °C for 1-3 h and hydrogenation in H₂ at 450 °C for 3 h were conducted (Table 3.3). Even for the nitrogenation at 300 °C, NdFe₇ absorbed nitrogen to form the corresponding metal nitride NdFe₇N_{1.88}. The nitrogen content decreased down to NdFe₇N_{0.89} after the subsequent hydrogenation made by the heat

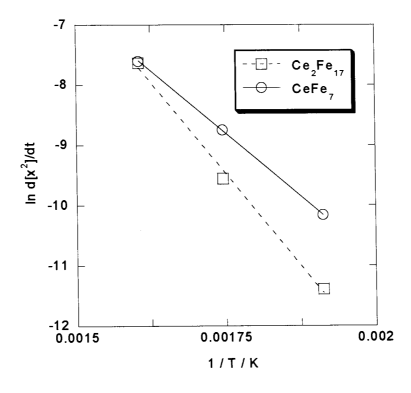


Fig. 3.4. Arrhenius plot $\rm s$ of the nitrogen desorption rate of CeFe₇ and Ce₂Fe₁₇.

treatment in H₂ at 450 °C. The Fe metal powder, however, was hardly nitrogenated under the same condition. This fact indicates that NdFe₇ is more effectively nitrogenated than Fe. By calculating based on the difference of nitrogen content (Δx =0.99) between NdFe₇N_{1.88} (after nitrogenation) and NdFe₇N_{0.89} (after hydrogenation), the quantity of the nitrogen desorbed from 1 cm³ of the metal nitride (ρ =ca.7.83 g cm⁻³) was converted to 14.46 mmol of NH₃, because the atomic nitrogen stored in the crystal lattice was released as NH₃ at about 100% of yield in the desorption step (see Fig. 3.2).

Table 3.3. Nitrogen absorption and desorption characteristics for the NdFe₇ and Fe

`	•	•		•	
Metal nitride (M.N.)	Cycle	Nitrogen content (x)		Regenerated ammonia	
	number	Nitrogenated	Hydrogenated	(mmol cm ⁻³ of M.N.) ^{b)}	
	(A) Nitrogenation in	NH₃ at 300 °Cfor 3	3 h	
$NdFe_7N_x$	1	1.88	0.89	14.46	
FeN_x	1	0	0	0	
	(B) Nitrogenation in	NH₃ at 450 °Cfor	1 h	
$NdFe_7N_x^{a)}$	1	3.90	1.93	28.83	
FeN_x	1	0.18	0.001	25.26	
	(C) Nitrogenation in	$\rm NH_3$ at 450 $^{\circ}\rm Cfor$	3 h	
NdFe ₇ N _x ^{a)}	1	4.08	1.81	33.21	
	10	3.43	0.83	38.03	
FeN_x	1	0.27	0.001	37.67	

a) Nitrogen contents x of the RFe₇N_x samples are represented as the values per chemical formula unit. b) The amount of ammonia regenerated is calculated by assuming that the nitrogen released from 1 cm³ of metal nitride is converted to ammonia.

The nitrogenation at 450 °C allowed the RFe₇ crystal lattices to absorb the much more amounts of nitrogen than that at 300 °C, owing to the enhanced nitrogen diffusion rate. As described above, however, the crystal lattices of RFe₇ were amorphized and NdFe₇ partly decomposed to precipitate α-Fe during the nitrogen absorption and desorption for them. Thus, such large amounts of nitrogen should be understood to be

stored in the composite materials, Fe/a-(Nd, Fe)N_x. Amounts of the ammonia regenerated from the Fe/a-(Nd, Fe)N_x composite powders were evaluated to be 33.21 mmol per 1 cm³ of the metal nitrides on the basis of the nitrogen analysis data for the nitrogenated and hydrogenated NdFe₇ one. It is noted that the amounts of the nitrogen regenerated from 1 \mbox{cm}^3 of Fe/a-(Nd, Fe)N $_x$ are 2.8 times larger than those of the conventional high-pressure nitrogen containers charged at 15 MPa (6.12 mmol of N₂ gas per 1 cm³ of their containers' Nd metal used as the starting materials were inactive for the dissociation of NH₃ at 450 °C to yield no metal nitride, and only a trace of NdN was observed during the nitrogenation at 700 °C. The Fe powder (particle size about 50 µm) also hardly reacts with NH₃ at 300 °C, whereas the RFe₇ intermetallic compounds can incorporate large amounts of nitrogen into their lattices. By raising the reaction temperature, however, a considerable amount of nitrogen was absorbed in the Fe crystal lattice. Consequently, the regenerated ammonia amounted to 37.6 mmol per 1 cm³ of FeN_x, although the formation of iron nitrides was required the higher reaction temperature compared with that of RFe₇. The XRD pattern of nitrogenated Fe powder was mixed with those of Fe₃N and Fe₄N. The amount of the ammonia regenerated for the Fe powder was comparable to that for the RFe₇ one in 10th cycle. In the case of nitrogenation for 1 h, the resulting quantity of ammonia for NdFe₇ is superior to that for Fe. The RFe₇ intermetallic compounds can absorb nitrogen faster than Fe under the more moderate reaction conditions.

The nitrogen storage capacity of NdFe₇ compounds increased with repeating the nitrogen absorption-desorption cycle. The amount of the ammonia regenerated from the Fe/a-(R, Fe)N_x composite powder in the 10th cycle (38.03 mmol) was higher than that in the first cycle (33.21 mmol). The XRD pattern of composite material after 10th nitrogenation was mixed with those of Fe₃N and a-(Nd, Fe)N_y. This means that these kinds of composite materials, Fe₃N/a-(R, Fe)N_y, give such excellent nitrogen absorption and desorption characteristics. It is concluded that the FeN_x/a-(R, Fe)N_y composite material is also highly active for the dissociation of NH₃. In the study of ammonia

synthesis over rare earth intermetallic compounds by Takeshita *et al.* [28], it has been reported that these intermetallic compounds decompose to α -Fe and RN during the reaction of N₂-H₂ mixed gas under a pressure of 7 MPa and the resulting α -Fe/RN composite materials are highly active for the production of ammonia.

Figure 3.5 shows scanning electron microscope photographs of the as obtained, 1st, and 10th nitrogenated NdFe₇ powders. As seen in Fig. 3.5a, it was found that many cracks occurred on the surface of the NdFe₇ particles because of the lattice expansion by absorbing large size of nitrogen atom interstitially. After the 10th nitrogenation cycle (see Fig. 3.5b), the particle size became smaller than the 1st one due to the lattice strain by nitrogen absorption and the phase separation took place between FeN_x and a-(Nd, Fe)N_y much more. The tendency that the amount of the ammonia regenerated from the metal nitride increases in 10th cycle is ascribed to the enlargement of the surface area since the crack formation becomes so frequent on the surface of the RFe₇ particles *via* the expansion and shrinkage of the lattices induced by repeating the nitrogen

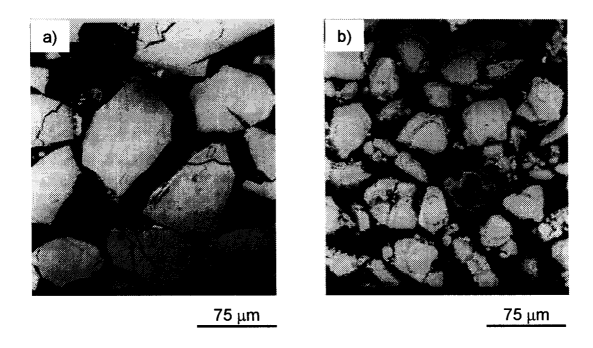


Fig. 3.5. SEM photographs of $NdFe_7N_x$ and the derived composite materials: a) 1st and b) 10th nitrogenated in NH_3 .

absorption-desorption cycle and the high activity for the dissociation of NH₃ molecules over the Fe₃N/a-(Nd, Fe)N_y composite materials.

Figure 3.6 shows the decomposition characteristics of ammonia over the NdFe₇ and Fe powders. The temperature to start the dissociation of ammonia over the NdFe₇ powder was lower than that over the Fe powder. This result indicates that the activity for the dissociation of ammonia over the NdFe₇ powder is higher than that of the Fe powder. The ammonia dissociation characteristics over the NdFe₇ and Fe powders fairly agreed with their nitrogen absorption properties as summarized in Table 3.3. Therefore, the nitrogenation over the RFe₇ compounds is expected to take place faster than Fe, so that ammonia is reproduced by cycling the nitrogenation and hydrogenation processes alternately in nitrogenation even at 300 °C.

As mentioned above, the RFe₇ intermetallic compounds can reversibly absorb and desorb significant amounts of nitrogen, and the nitrogen incorporated into the crystal lattices is easily converted to ammonia in the nitrogen desorption (hydrogenation) step.

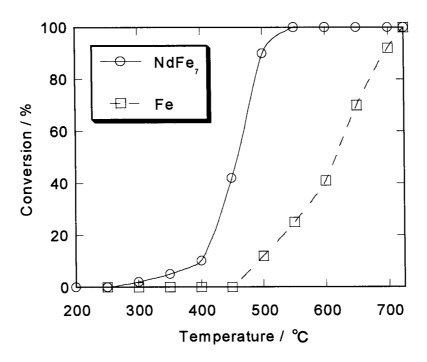


Fig. 3.6. Decomposition characteristics of ammonia over the CeFe₇ and Fe powders.

However, this nitrogen absorption-desorption system is less important in the viewpoint of the chemical engineering for use of ammonia as nitrogenation source. If molecular nitrogen is used as such nitrogen source, it is possible to develop novel reaction process using atomic nitrogen by which molecular nitrogen can be eventually converted to ammonia via the nitrogenation-hydrogenation cycle for the intermetallic compounds. For the catalytic approaches, the ammonia synthesis has been already tried over the rare earth-transition metal intermetallic compounds (e.g. CeRu₂, HoFe₃, and PrCo₅) by Takeshita and coworkers [28]. After conducting the reaction, the crystal structure of HoFe₃ completely decomposed to HoN and α-Fe. This result suggests that HoN hardly releases nitrogen owing to its great stability of ionic bonding-type nitride and the ammonia formation by the direct reaction between N₂ and H₂ gas (under a high pressure condition of about 7MPa) mainly takes place on the surface of transition metal promoted by electron transferring from rare earth nitrides such as HoN. In this study, the nitrogenation and hydrogenation processes are performed alternately in the individual N₂ and H₂ streams, and consequently, the atomic nitrogen desorbed from the metal nitride is converted to ammonia in the desorption step under atmospheric pressure.

The nitrogenation for CeFe₇ was conducted in N_2 instead of NH₃ by heating at 673 K for 3 h. The nitrogen content per chemical formula was 0.70. The subsequent nitrogen desorption was carried out in H₂ at 723 K for 3 h to desorb nitrogen down to CeFe₇N_{0.66}. The amount of ammonia product per unit volume was only 0.58 mmol that sharply decreased compared with the case of nitrogenation in NH₃. The respective binding energy of N-H and $N \equiv N$ are 431 and 941 kJ mol⁻¹. The dissociation of strong triple bonding of N_2 is assumed to be rate determining step for nitrogen absorption reaction. It is well known that ammonia is more favorable as the nitrogen source than molecular nitrogen in the points of thermodynamics [52]. To accelerate the dissociation ability for molecular nitrogen, the composite material between the Ru/La₂O₃ catalyst (15 wt. %) and the CeFe₇ intermetallic compound (85 wt. %) was prepared by the mechanical

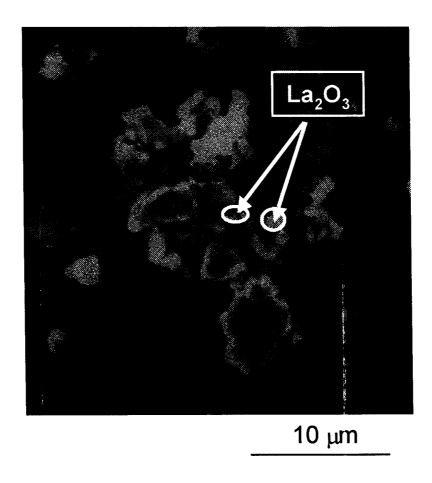


Fig. 3.7. Scanning electron microscope photograph of the Ru/La_2O_3 -CeFe $_7$ composite powder.

grinding method, of which the Ru/La₂O₃ catalyst plays a role of promoting the dissociation of molecular nitrogen reported by Aika et al [46]. From the scanning microscope measurement with an EPMA analysis, it was confirmed that the La₂O₃ was loaded on the surface of CeFe₇ (see Fig. 3.7).

For the Ru-La₂O₃/CeFe₇ composite powder, the nitrogenation was performed at 673K in N₂ for 3 h. The amount of nitrogen absorption per chemical formula increased to be 1.56. After the subsequent nitrogen desorption by heating at 473K in H₂ for 3 h, the nitrogen content per chemical formula unit decreased to 1.38, resulting that the amount of ammonia product per unit volume was up to 2.91 mmol. At 5th cycle for nitrogen absorption and desorption over Ru-La₂O₃/CeFe₇, the nitrogen content per

chemical formula unit after nitrogenation was 1.65, whereas 1.43 after denitrogenation. The amount of ammonia product per unit volume was consequently 3.55 mmol (Table 3.4). Ru-La₂O₃/CeFe₇ showed a good durability for nitrogen absorption-desorption cycle. The resulting amount of ammonia product per 1 g of CeFe₇ at 5th cycle is 452 µmol. From these results, the Ru-La₂O₃/CeFe₇ composite material can store highly reactive atomic one in the crystal lattice with a capacity of 40 times volume of nitrogen per its unit volume. Since the highly reactive monoatomic nitrogen easily reacts with

Table 3.4 Nitrogen absorption and desorption characteristics for CeFe₇ and Ru-La₂O₃/CeFe₇

Metal nitride	Cycle number	Nitrogen content (x) ^a		Regenerated
		After nitrogenation	After hydrogenation	ammonia (mmol/cm ³)
CeFe ₇ N _x	1	1.31	1.21	1.6
Ru-La₂O₃ /CeFe ₇ N _x	1	1.56	1.38	2.91
	5	1.65	1.43	3.55

a) Nitrogen contents are represented as the values per chemical formula unit.

hydrogen to generate ammonia at about 100 % of yield, it is concluded that the above nitrogen absorption-desorption reactions is regarded as a new process of ammonia synthesis using the formation of interstitial type metal nitrides.

3.4 Conclusions

Rare earth-iron intermetallic compounds RFe₇ (R=Ce, Nd and Sm) reversibly absorb and desorb large amounts of nitrogen by cycling the respective nitrogenation and hydrogenation processes made in NH₃ and H₂ at 300-450 °C. The amounts of the nitrogen regenerated per unit volume of these compounds through the nitrogen absorption and desorption is larger than that of the conventional high pressure nitrogen cylinder charged

at 15 MPa. The nitrogen storage capacity of the RFe₇ compounds are superior to that of the R₂Fe₁₇ ones since the crystal structure of RFe₇ is the iron deficient Th₂Zn₁₇-type one which is favorable for the nitrogen diffusion in the bulk. Additionally, the atomic nitrogen stored as the metal nitrides is regenerated as ammonia in the nitrogen desorption step in H₂ at about 100% of yield. The amount of the regenerated ammonia is increased with repeating the nitrogen absorption-desorption cycle of RFe₇ compounds and the formation of FeN_x/a-(R, Fe)N_y composite materials is responsible for the increase of nitrogen regeneration. Futhermore, RFe₇ can absorb nitrogen by the nitrogenation in N₂ at 400 °C even though molecular nitrogen is generally too hard to produce such metal nitrides because of the huge bonding energy. The resulting nitrides RFe₇N_x release the nitrogen as ammonia by heating in H₂. The Ru/La₂O₃-CeFe₇ composite powder was more easily nitrogenated by N₂ in order to promote the ability for dissociation of molecular nitrogen by Ru/La₂O₃ catalyst, resulting that the larger amounts of ammonia generation was than CeFe₇. Ru/La₂O₃-CeFe₇ can generate larger amount of ammonia compared with Ru-Al₂O₃-TiFe₂ even at lower temperature of nitrogenation in N₂.

Summary

Chapter 1.

The intermetallic compounds R₂Fe₁₇ (R=Y, Ce, and Sm) formed the corresponding interstitial type metal nitrides by heating in a mixed gas of NH₃-H₂ at 350-450 °C. The nitrogen stored as the metal nitrides was reversibly desorbed as NH₃ at about 100% of yield by the following heating in H₂ at 450 °C. The amount of the nitrogen released per unit volume of these intermetallic compounds is larger than that of a conventional nitrogen container charged at 15MPa. The nitrogen storage capacity of Sm₂Fe₁₇N_x increased by repeating the nitrogenation-hydrogenation cycles owing to the formation of FeN_x/a-(R, Fe)N_y composites with large surface areas derived from the starting intermetallic compound through nitrogenation and hydrogenation. Additionally, the nitrogen storage property of Sm₂Fe₁₇ powder was effectively improved by surface modification with Ru metal loading.

Chapter 2.

Nitrogen absorption and desorption characteristics were conducted for the intermetallic compounds such as MFe₂ (M=Ti, Zr, Nb, and Mo) or M'Fe (M'=Ti and V) Among them, a series of the Laves-phase MFe₂ and σ-phase VFe compounds absorbed large amounts of nitrogen into the crystal lattices to transform to the amorphous-like metal nitrides, of which the randomized structures were induced by the insertion of large atomic size of nitrogen into the crystal lattices. These metal nitrides reversibly released the nitrogen by the following hydrogenation, and the amorphous-like structures were maintained after that. Furthermore, once such amorphous-like modification of TiFe₂ (a-TiFe₂) induced by the nitrogenation in the NH₃-H₂ mixed gas could absorb nitrogen even by the nitrogenation in N₂ gas with co-loaded with the ruthenium metal and aluminum oxide on the surface of TiFe₂. The nitrogen stored was also released as

ammonia in the subsequent hydrogenation under atmospheric pressure.

Chapter 3.

Reversible nitrogen absorption and desorption were observed on the rare earth-iron intermetallic compounds RFe₂ and RFe₇. In particular, the intermetallic compounds RFe₇ with the modified Th₂Zn₁₇-type structure absorbed large amounts of nitrogen into their crystal lattices. The nitrogen absorption and desorption properties over Th₂Zn₁₇-type RFe₇ were superior to those over Th₂Zn₁₇-type R₂Fe₁₇ due to the ease of nitrogen diffusion in the bulk, since Fe (6c) site is partially occupied (occ=2/3) in the crystallographic cell for RFe₇, whereas Fe (6c) site is fully occupied (occ=1) for R₂Fe₁₇. Furthermore, CeFe₇ could absorb nitrogen even by nitrogenation in N₂ at 400 °C. The resulting metal nitride desorbed nitrogen as ammonia by subsequent. Ru-La₂O₃/CeFe₇ composite powder, which was active for dissociation for molecular nitrogen by loading Ru-La₂O₃ catalyst, can product more larger amount of ammonia than Ru-Al₂O₃/a-TiFe₂ even at lower nitrogenation temperature *via* the nitrogenation-hydrogenation cycle.

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