TITILE: Thermal and Structural Characterization of Nanocomposite Iron Nitride - Alumina and Iron Nitride - Silica Particles

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Thermal and Structural Characterization of Nanocomposite Iron Nitride - Alumina and Iron Nitride - Silica Particles

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ABSTRACT

Nanocomposite iron nitride based powders are known to have enhanced magnetic and other physical properties. To further explore their potential for application in various fields, we have performed a systematic study of the iron nitride - alumina and iron nitride - silica systems. Iron nitride powder of composition FeₙN (2 < x < 4), containing both Fe₃N and Fe₄N phases, was mechanically milled with Al₂O₃ or SiO₂ powder for 4, 8, 16, 32, and 64 hours at the following compositions: 
(FeₓN)₀.₂(Al₂O₃)₀.₈, (FeₓN)₀.₆(Al₂O₃)₀.₄, (FeₓN)₀.₂(SiO₂)₀.₈, and (FeₓN)₀.₆(SiO₂)₀.₄. Differential thermal analysis and X-ray diffraction were performed to investigate thermal and structural transitions as a function of milling time. As the milling time is increased, the thermal peak corresponding to Fe₄N is diminished, while the one corresponding to Fe₃N is enhanced. These transitions are correlated with X-ray diffraction patterns. All XRD peaks broaden as a function of milling time, corresponding to smaller particle size. Transmission electron microscopy also reveals a decrease in particle size as the milling time in increased.

INTRODUCTION

Small magnetic particles are important in many practical applications for their superior magnetic properties such as high saturation magnetization and high coercive field. Improvements in these properties have been obtained through sputtering [1,2] and mechanical alloying [3] of iron particles with alumina or silica, an improvement which has been correlated with a reduction in particle size. Metal-metalloids, such as FeₙN, have these desirable magnetic properties, but conventional techniques used to produce small particles often lead to oxidation and the formation of an iron oxynitride surface layer, which is ferromagnetically coupled to the core of a particle. Surface oxidation may be avoidable by embedding single domain FeₙN particles in an insulating matrix such as Al₂O₃ or SiO₂. In fact, iron particles dispersed in such an insulating matrix have shown enhanced magnetic and mechanical properties [3-5]. In this work, small FeₙN particles are embedded in an insulating matrix through mechanical ball-milling. The resulting particles should have the advantageous properties of both small particle size and metal-metalloid composition. To fully understand the mechanisms producing these particles, structural and thermal studies are performed in this study to evaluate the phase formation as a function of milling time.
EXPERIMENTAL DETAILS

Samples of iron nitride - alumina or iron nitride - silica were formed by mechanically milling powder of Fe₃N (2<x<4) with Al₂O₃ or SiO₂ powder for 4, 8, 16, 32, or 48 hours in a Fritsch planetary ball mill. The 325 mesh powders were 99.9% pure. Oxidation during milling was prevented by milling in the presence of ethyl alcohol. The mixture of Fe₃N powder to Al₂O₃ or SiO₂ powder was varied so that samples at each milling time were obtained for (Fe₃N)ₓ(Al₂O₃)₁₋ₓ and (Fe₃N)ₓ(SiO₂)₁₋ₓ with y = 0.2 and 0.8 (z = 0.8 and 0.2, respectively).

Thermal analysis was performed using an Instrument Specialists differential thermal analyzer with Pt/PtRh thermocouples and alumina sample and reference cups. Al₂O₃ powder was used as the reference material and scans were carried out with a heating / cooling rate of 10°/minute. Powder X-ray diffraction data was obtained with a Philips PC-APD3520 diffractometer using Cu Kα radiation. Samples were prepared for transmission electron microscopy (TEM) by sonicating the powder in methanol and then placing a few drops of the solution onto a holey-carbon coated grid. TEM was performed on a JEOL 1200 EX. TEM negatives were scanned at 1200dpi and then analyzed using the National Institute of Health's (NIH) Image software to measure particle size.

RESULTS AND DISCUSSION

Figure 1 shows XRD plots for the milled compounds with y = 0.2 (z = 0.8) and the pure Fe₃N powder. The starting Fe₃N powder (2 < x < 4) is a mixture of Fe₃N (ε phase) and Fe₅N (γ').

Figure 1: XRD patterns for (a) (Fe₃N)₀.₂(Al₂O₃)₀.₈ and (b) (Fe₃N)₀.₂(SiO₂)₀.₈ for the various milling times and for the starting Fe₃N powder.
The peaks at 44.5° and 65.0° could be either α-Fe or γ'-Fe₄N. For the Al₂O₃ series (figure 1a), it is clear that peaks corresponding to Fe₄N decrease in intensity with milling time. In particular, the Fe₄N peaks at 48° and 70° completely disappear at even short milling times. An increase in milling time also correlates with an increase in the intensity of the Fe₃N peaks, particularly those at 43.3° and 69°. This suggests that milling induces a transition from Fe₄N to Fe₃N. Thus, there is an excess of Fe, which is evidenced by the enhancement of α-Fe peaks at 44.5° and 65°. The peak at 41° (γ' and ε) grows on high side with milling time, which indexes more to the Fe₃N phase. A similar trend is seen for the SiO₂ series (figure 1b). The Fe₃N peaks increase in intensity while the peaks corresponding to the γ' phase decrease. Again, the persistence of the peak at 44° suggests a growth of α-Fe due to excess Fe coming from the Fe₄N to Fe₃N transition.

Similar results were obtained for the higher Fe composition samples; (FeₓN)₀.₆(Al₂O₃)₀.₄ and (FeₓN)₀.₆(SiO₂)₀.₄. All of the XRD peaks broaden as milling time increases, indicates a decrease in particle size.

Figure 2 displays the DTA traces for both series as well as the curve for the starting Fe₄N powder. The ε phase (Fe₃N) has a wide range of solubility and its decomposition temperature at 25% N (atomic percent) occurs over a range of temperatures, but is less than 680°C for bulk material. Fe₄N, with a much narrower range of solubility, decomposes at 680°C for bulk material [6]. For small particles, these decompositions occur at lower temperatures. Decomposition temperatures of 570°C have been reported for 10μm Fe₄N particles [7], and 410°C-582°C has been reported for 0.3-5μm Fe₃N particles [8]. Table I displays our average

![Figure 2: DTA curves for (a) the (FeₓN)₀.₂(Al₂O₃)₀.₈ series and (b) the (FeₓN)₀.₆(Al₂O₃)₀.₄ series. The y-axis is ΔT in °C, but normalized to show multiple curves on one graph. The peaks shown covered a range in ΔT of 20°C.](image)
Table I  Average particle size as measured by TEM studies for the Al₂O₃ samples.

<table>
<thead>
<tr>
<th>Milling Time (hr)</th>
<th>Average Particle Size (nm)</th>
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<tbody>
<tr>
<td></td>
<td>(Fe,N)₁₋₂₃(Al₂O₃)₀ₓ</td>
</tr>
<tr>
<td>4</td>
<td>376</td>
</tr>
<tr>
<td>8</td>
<td>278</td>
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<tr>
<td>16</td>
<td>172</td>
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<tr>
<td>32</td>
<td>159</td>
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<tr>
<td>64</td>
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particle sizes for the Al₂O₃ samples as measured by TEM. Since our particle sizes are all less than 0.5µm, the decomposition of the nitride phases will occur at lower temperatures. For the starting powder, which is a mixture of Fe₃N (ε) and Fe₄N (γ), two exothermic peaks appear in range for nitride decomposition (400-600°C). The prominent peak at 425°C likely represents Fe₃N decomposition, and the peak nested in its high temperature shoulder represents the decomposition of Fe₄N.

For the Al₂O₃ samples, the Fe₃N peak becomes more prominent with milling time. This peak also shifts to slightly lower temperature, which correlates with the decrease in particle size upon increased milling time. This shows, as did the XRD data, the Fe₃N being converted to Fe₄N upon increased milling time. The DTA curves also show the growth of a peak between 300°C and 350°C as milling time increases. If the excess Fe is going into the α phase, as suggested by the XRD, this peak could correspond to a transition from γ to α-Fe as N is liberated. Both of these phases have a wide range of solubility for N below 590°C. These lowest temperature peaks could also be due to adsorbed water or the presence of hydrocarbons in the samples, which were milled in ethyl alcohol to prevent oxidation. The two nitride phase peaks are most prominent in the samples with lower milling time (see the 4, 8, 16hr samples for (Fe₃N)₀₋₆ₓ(Al₂O₃)₀₋₄ₓ, figure 2b). As the milling time increases, the lower temperature peak is more prominent. The shift to lower temperature for the Fe₃N peak is also most prominent in the (Fe₃N)₀₋₆ₓ(Al₂O₃)₀₋₄ₓ series, although these particles are larger on average than those in the (Fe₄₄₋₈ₓ(Al₂O₃)₀₋₈ₓ samples. The higher concentration of Al₂O₃ is therefore partly responsible for the decomposition of the nitride phases.

A similar trend is seen for the SiO₂ samples (figure 3). For the (Fe₃N)₀₋₂ₓ(SiO₂)₀₋₈ (figure 3a) series, it is clear that the peak corresponding to Fe₃N (400-450°C) grows with increasing milling time, indicating the presence of more Fe₃N phase. This trend is also seen for the (Fe₄₄₋₈ₓ(SiO₂)₀₋₄ series (figure 3b), with the exception of the 64hr and perhaps the 4hr sample. In these two cases, the Fe₄N peak is more prominent. The three prominent peaks between 200°C and 375°C for the 64hr sample suggest that it may have oxidized or otherwise been contaminated. More studies should be performed to verify this suggestion.

CONCLUSIONS

We have explored ball milling of Fe₃N and an insulating component (alumina or silica) as a means of producing ultrafine Fe₃N particles. A reduction of the Fe₃N particle size is seen as
Figure 3: DTA curves for (a) the \(\text{(Fe,N)}_0.2\text{(SiO}_2\text{)}_{0.8}\) series and (b) the \(\text{(Fe,N)}_{0.4}\text{(SiO}_2\text{)}_{0.6}\) series. The y-axis is \(\Delta T\) in °C, but normalized to show multiple curves on one graph. The peaks shown covered a range in \(\Delta T\) of 20°C.

Milling time increases, and this is accompanied by a transition from Fe₄N to Fe₃N. Both structural and thermal characterization show that there is an increase of the \(\varepsilon\) Fe₂N phase as milling time is increased. This decrease in size causes the decomposition of the Fe₃N phase to shift to a lower temperature, and a broadening of lines in the XRD spectra. Studies of the magnetic properties of these ultrafine particles have been completed [9] and suggest that ball milling iron nitride particles is a viable technique to produce small particles with desirable magnetic properties.

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REFERENCES