PREPARATION OF Fe/GLASS COMPOSITE BY REDUCTION OF Na₂O–Fe₂O₃–B₂O₃–SiO₂–ZnO GLASS AND GLASS CERAMICS

A. Mohammadpour, S. M. Mirkaemi and A. Beilollahi

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School of Metallurgy and Materials Engineering, Iran University of Science and Technology, Tehran, Iran.

Abstract: In the present study, the feasibility of α-Fe ferromagnetic phase formation in glass and glass-ceramic by reduction in hydrogen atmosphere have been investigated. The glass with the composition of 35Na₂O–24Fe₂O₃–20B₂O₃–20SiO₂–1ZnO (mol %) was melted and quenched by using a twin roller technique. As quenched glass flakes were heat treated in the range of 400–675 °C for 1–2 h in hydrogen atmosphere, which resulted in reduction of iron cations to α-Fe and FeO. The reduction of iron cations in glass was not completely occurred. Saturation magnetization of these samples was 8.37 emu g⁻¹. For the formation of glass ceramic, as quenched glass flakes heat treated at 590°C for 1 h. Heat treatment of glass ceramic containing magnetite at 675°C in hydrogen atmosphere for 1 h led to reduction of almost all of the iron cations to α-Fe. Saturation magnetization of this sample increased from 19.8 emu g⁻¹ for glass ceramic to 67 emu g⁻¹.

Keywords: Magnetite, α-Fe, Reduction, Glass, Glass Ceramic, Magnetic Composite

1. INTRODUCTION

Soft magnetic composites (SMCs) are ferromagnetic particles coated with a thin electrically insulating layer and then pressed into the desired shape by the powder metallurgy (PM) methods. SMCs have unique magnetic properties, such as a three-dimensional (3D) isotropic ferromagnetic behavior, low eddy current loss, relatively lower total core loss at medium and high frequencies, and flexible design, which can be applied to various fields [1–5]. To provide the maximum magnetic permeability, the amount of inter-particle insulation should be minimized and the iron content maximized. Several researchers have tried to improve the magnetic performance of SMCs by selecting suitable insulating materials and applying various coating methods [6, 7]. Currently, insulating coatings include epoxy, polyamide, silicon resins and polyvinyl alcohol [8–10]. The thermal treatment temperature for these materials is limited by the thermal resistance of the organic insulating layer between the magnetic particles; therefore phosphates (zinc/iron/manganese) and oxides (MgO, SiO₂) with high resistance temperatures were selected instead of organic coatings [11–12]. Various methods have been explored to synthesize metal/insulator nanocomposites, including sol–gel [13], sputtering [14], electrodeposition [15], high energy ball milling [16], microemulsion and reverse micelle techniques [17]. Most of these methods are complex and expensive with low yield. In the present study, a new kind of SMC with glass matrix was obtained by a simple method which was reduction of glass and glass ceramic containing Fe cations. This novel method could be used to produce SMCs with different compositions and magnetic properties with low cost.

2. EXPERIMENTAL

2.1. Glass Formation

The nominal chemical compositions of prepared glasses was 35Na₂O–24Fe₂O₃–20B₂O₃–20SiO₂–1ZnO (mol %). The raw materials were reagent grade Na₂CO₃, CaCO₃, Fe₂O₃, ZnO, SiO₂, and B(OH)₃. The batches were melted in alumina crucibles in 1300–1350 °C range, with a holding time of 15 min at the final temperature. The melts were then quenched between rotating steel rollers.
2. 2. Glass-Ceramics Formation

For crystallization of glass, resulting flakes ~50 µm thick were heat treated under a graphite powder bed. According to previous research [18] time and temperature of heat treatment was selected 1 h and 590 °C respectively.

2. 3. Reduction of Glass and Glass Ceramics

Flakes of glass and powder of glass ceramic (<53 µm) samples were introduced in a silica tube placed in a furnace. They were heated under argon flow until the desired treatment temperature was reached. Then argon was replaced by hydrogen (450-500 ml/min) at atmospheric pressure. Treatment times were between 1 and 2 h, and temperature in the range 400 to 675 °C. At the end of this time, hydrogen was replaced by argon and the furnace was allowed to cool to room temperature by switching off the power. One of the glass samples was maintained in 590 °C for 2 h in Ar atmosphere before reduction. The synthesis condition of samples is shown in table 1.

2. 4. Characterization

The phase identification of the samples were studied by X-ray diffraction (XRD) (PANalytical X´Pert Pro MPD) using Cu Kα radiation. The magnetization measurements were carried out by vibrating sample magnetometer (VSM) MDK6 Meghnatis daghigh kavir, mizan daghigh.

3. RESULT AND DISCUSSION

3. 1. Glass and Glass Ceramic

XRD patterns of the glass (Fig. 1) illustrates that the sample is amorphous in the accuracy range of XRD analysis.

Fig. 2 depicts the XRD pattern of glass ceramic. It reveals the existence of α-Na₂Si₃O₅ and α-Na₂B₂O₅ phases besides magnetite in this sample. Using Scherrer's equation the mean particle size of magnetite phase in this sample was calculated as 21 nm. Also The X-ray diffraction pattern of the sample contains the peaks of iron silicate phases. By consuming Fe cations, the formation of these phases leads to a decrease in the amount of magnetite phase in final glass ceramic.

Hysteresis loop of glass and glass ceramic is shown in fig. 3 and 4 respectively. Maximum magnetization of the glass was 1.6 emu g⁻¹ and the saturation wasn’t observed in the curve. The value of the magnetization is probably due to the formation of magnetite phase during cooling of the glass melt and it is not detectable by XRD analysis because the amount of crystalline phase was negligible. The reason that curve has not been reached to the saturation could be the possible existence of superparamagnetic particles in glass [18].

The saturation magnetization (Ms) of the glass

<table>
<thead>
<tr>
<th>Sample</th>
<th>Crystallization</th>
<th>Reduction in H₂ atmosphere</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Temp. (°C)</td>
<td>Time (h)</td>
</tr>
<tr>
<td></td>
<td>Temp. (°C)</td>
<td>Time (h)</td>
</tr>
<tr>
<td>GC</td>
<td>590</td>
<td>1</td>
</tr>
<tr>
<td>GR-400-2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>CR-590-1</td>
<td>400</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>GR-657-2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>GR-657-2</td>
<td>675</td>
<td>-</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>GCR-675-1</td>
<td>590</td>
<td>1</td>
</tr>
<tr>
<td>675</td>
<td>1</td>
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</tr>
</tbody>
</table>

Fig. 1. XRD pattern of glass (G).
Fig. 2. XRD pattern of glass ceramic (sample GC- 590°C, 1h).

ceramic (sample GC) is 19.8 emu g⁻¹ and coercive field (Hc) is negligible. The value of the maximum Ms obtained for glass ceramic is different from the theoretically calculated value. If the maximum amount of theoretically feasible magnetite was crystallized in the GC sample, it would have a magnetization value of ~40 emu g⁻¹. This value is obtained by multiplication of the theoretically possible maximum amount of magnetite in this sample GC (~43 wt.%) by the magnetite theoretical magnetization value (92 emu g⁻¹ [19]).

One of the reasons for this difference could be related to incomplete growth of the magnetite

Fig. 3. Hysteresis loop of sample glass.
crystals and their small crystallite size. The saturation magnetization of the magnetite nanoparticles with particle sizes about 7-13 nm have been reported in the range of 52 – 75 emu g⁻¹ [20].

As observed in the X-ray diffraction pattern of glass ceramic, FeSiO₃ and NaFe(SiO₃)₂ iron silicate phases were formed during the heat treatment. These phases are non-magnetic and by consuming some of the iron cations in the glass, it would be feasible to decrease the amount of crystallized magnetite phase. Therefore the saturation magnetization would be decreased in the glass ceramic.

3. 2. Reduction of Glass and Glass Ceramic

Fig. 5 shows the glass flakes heat treated at 400 °C for 2 h in hydrogen atmosphere (sample GR-400-2). The peaks corresponding to magnetite was observable in the X-ray diffraction pattern. These relatively broad peaks represent of the existence of a fine crystallite size of this phase. Also, from the high background in the X-ray diffraction pattern of the sample could be understood that the amount of crystallized phase in this sample would be very low. The characteristic peak of α-Fe in the X-ray diffraction pattern wasn’t observed, so the reduction of glass in 400 °C for 2 h didn’t lead to the formation of α-Fe phase. The reason could be the low permeability and diffusivity of hydrogen in glass at the reduction temperature.

Fig. 6 shows the XRD pattern of the glass flakes heat treated at 590 °C for 1 h in hydrogen
atmosphere (Sample GR-590-1). It can be seen that by increasing the heat treatment temperature, the α-Fe and wustite phases have been crystallized with magnetite phase in H₂ atmosphere. The existence of iron phase shows the reduction of iron cations to metallic iron. The hydrogen gas was individually applied in order to reduce glass and the heat treatment temperature was much higher than the Tg of the glass (373 K) [18]. As a result, the high amount of hydrogen permeability and solubility in glass was expected. It is reported that the reduction mechanism of cations in glass is controlled by the permeability of hydrogen in high hydrogen partial pressure (over 10%) atmosphere and at heat treatment temperatures higher than Tg of glass [21]. Iron cations could have been reduced in two ways. First, magnetite phase was crystallized in glass and then it reduced to α-Fe. In the other way, the overall reduction process could be the result of hydrogen diffusion to the site of the iron ions, reduction of the iron ions to the atomic state, diffusion of these atoms to a nuclei, and subsequent growth of these nuclei to form particles of α-Fe [22].

The existence of wustite in the glass heat treated in H₂ atmosphere could be due to the reduction of Fe³⁺ to Fe²⁺ cations. Also, it has been reported that at temperatures higher than 570 °C, the reduction of magnetite proceeds through the formation of non-stoichiometric wustite [23]. It seems that crystallized magnetite phase reduced during heat-treatment in hydrogen atmosphere and the existence of wustite was probably due to the incomplete reduction of the magnetite phase for 1 hour.

In comparison with sample GR-590-1, in sample GR-675-2, the time and the temperature of the heat treatment in hydrogen atmosphere was increased in order to enhance the hydrogen permeability in glass which would result in reduction of magnetite to the α-Fe phase.

Fig. 7 shows X-ray diffraction pattern of glass flakes reduced in hydrogen atmosphere for 2 h at 675 °C (GR-675-2). Crystallized phases in the sample GR-675-2 are observed in Fig 7. Besides peaks of α-Fe phase, the peaks corresponding to the magnetite (Fe₃O₄) and wustite (FeO) phases were detectable. In this sample, magnetite phase has not been completely transformed to α-Fe.

Before reduction of the sample GR-590-2-675-

![XRD Pattern](image)

**Fig. 6.** XRD pattern of GR-590-1 (590 °C, 1 h, H₂ atmosphere).

The existence of wustite in the glass heat 2, it was maintained at 590 °C for 2h in argon
Fig. 7. XRD pattern of GR-675-2 (heat treated at 675 °C for 2 h in H₂ atmosphere).

Fig. 8. XRD pattern of GR-590-2-675-2 (heat treated at 590 °C for 1 h in Ar atmosphere and at 675 °C for 2 h in H₂ atmosphere).
atmosphere for crystallization of the magnetite phase in glass. The crystallization and the reduction magnetite were carried out consecutively. XRD pattern of the sample GR-590-2-675-2 is shown in Fig. 8. As seen in this sample, the proportion of FeO peaks intensity to those of α-Fe increased compared to sample GR-675-2. Because FeO is an intermediate phase of reduction of magnetite to metallic iron, it was likely that the reduction process of magnetite was incomplete.

The powder of sample GC (<53 μm) was heat treated at 675 °C for 1 h in hydrogen atmosphere (GCR-675-2). Fig. 9 depicts the XRD pattern of the sample GCR-675-2. It is understood that in comparison to sample GC, the peaks of the magnetite were removed and the characteristic peaks of iron (α-Fe) were observed. Also iron containing silicate phases in the sample GC such as FeSiO₃ and NaFe₂(SiO₄)₂ were not detected in the XRD pattern of the sample GCR-675-1. This could be due to the reduction of the iron cations to α-Fe in these phases.

According to the magnetization results (Fig. 10 and 11) the sample Gr-400-2 containing a maximum magnetization value of 8 emu g⁻¹ and a very weak coercive force (which could not be measured). In this case, the magnetization never reaches the saturation level and there is no hysteresis loop as a result of presumably the small size of the magnetite particles, falling into the superparamagnetic range. In sample GR-590-1, the magnetization curve approached the saturation level and showed the Ms value of 27 emu g⁻¹. In this case the Hc value is 123 Oe. The Ms of GR-590-1 was measured higher than GC-590-1. This increase in Ms can be mainly attributed to the reduction of the magnetite to α-Fe phase with higher value of Ms than magnetite (α-Fe Ms=218 emu g⁻¹). Ms and Hc of sample GR-675-2 was estimated 37 emu g⁻¹ and 93 Oe respectively. By increasing time and temperature of the heat treatment in the sample GR-675-2 in comparison to the sample GR-590-1 probably the amount of the hydrogen permeability in glass increased and higher amount of cations has been reduced.

Hysteresis loop of a two-step reduced glass is given in Fig. 11. The saturation magnetization of this sample was measured 27 emu g⁻¹ and Hc was

![XRD pattern of GCR-675-1 (glass ceramic heat treated at 675 °C for 1 h in H₂ atmosphere).](image)

As observed, with the heat
treatment of the glass at crystallization temperature before the reduction, the saturation magnetization was reduced, presumably due to the incomplete reduction of magnetite and the increase in the FeO amount in the sample. Also some of the iron cations in glass consumed by some non-magnetic silicate phases which leads to the reduction of the overall magnetization. Direct reduction of glass flakes was unable to reduce all iron cations to metallic iron.

Hysteresis loop of reduced GC powder (GCR-675-1) is shown in Fig. 11. In this sample the magnetization was increased to 67 emu g⁻¹ which was about three times higher than that of GC. The
value of the maximum Ms obtained for GCR-675-1 is close to theoretically calculated value. If the maximum amount of theoretically feasible α-Fe was formed in the GCR-675-1 sample, it should have a magnetization value of ~77 emu g⁻¹. This value is calculated by multiplication of the theoretically feasible maximum amount of α-Fe in this sample GCR-675-1 (~36 wt.%) by the theoretical magnetization value of α-Fe (218 emu g⁻¹). The difference between Ms of sample GCR-675-1 and the calculated ones could be induced by changes in the composition during the heat treatment.

The ability of the formation of ferromagnetic phases in glass was observable. Saturation magnetization of ferromagnetic materials is higher than that of ferrimagnetics. Therefore, compared with glass ceramics, the total saturation magnetization of reduced glass ceramics was enhanced.

In the novel magnetic composite materials, obtaining various magnetic properties by changing in the ratio of magnetic phase, the microstructure and composition of ferromagnetic and glass phases might be possible.

4. CONCLUSION

By the use of the rapid roller quenching technique, glass flakes in the NaₓO–FeₓOₓ–B₂O₃–SiO₂–ZnO system were obtained. Heat treatment of the obtained glass at 590 °C for 1 h resulted in the crystallization of magnetite phase and the Ms value of 19.8 emu g⁻¹. Reduction of glass flakes at 590 °C for 1 h and 675 °C for 2 h led to the formation of the α-Fe phase with magnetite and wustite, which exhibited magnetization values of 27 and 37 emu g⁻¹, respectively. Reduction of almost all magnetite to α-Fe phase was observed in reduced glass ceramics containing magnetite phase in a hydrogen atmosphere at 675 °C for 1 h. Ms of this sample increased to 67 emu g⁻¹ which was three times higher than that of magnetic ceramic (19.8 emu g⁻¹). It seems that a new generation of glass/ferromagnetic composites can be achieved by the reduction of glass ceramics containing ferromagnetic metal compounds.

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