DEVELOPMENT OF HARDNESS IN A Fe- BASED NANO-COMPOSITE ALLOY

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Abstract: Achieving extreme hardness in the newly synthetic steel formed by converting from initial amorphous state to subse-quent crystalline structure -named as devitrification process- was studied in this research work. Results of TEM observations and XRD tests showed that crystallized microstructure were made up four different nano-scale phases i.e., α-Fe, Fe2Mo10Cr12, Fe5C and Fe3B. More, Vickers hardness testing revealed a maximum hardness of 18.6 GPa which is significantly harder than existing hardmetals. Detailed kinetic and structural studies have been proof that two key factors were contributed to achieve this extreme hardness: supersaturation of transition metal alloying elements (especially Nb) and also reduction in the structure to the nano-size crystals.

Keywords: Transmission electron microscopy (TEM); Nano composites; X-ray diffraction (XRD)

1. INTRODUCTION

"Devitrified nanostructured steels" achieving from initial glassy state have been attracted much attention during last decades because of their unique hardness [1]. The main approach to produce iron based nanocomposites is through solid/solid state transformation in which nanostructured steels are obtained from unstable initial amorphous structures by annealing process [2]. Indeed, this new class of steels was first developed by over-quenching from a metallic glass and then heat treating the glass precursor over its crystallization temperature to devitrify it to a multiphase nanocrystalline structure [3, 4].

It is worth noting that overquenched metallic glasses, existing in a supersaturated metastable condition, can be transformed to multi-phase crystalline structures when enough energy was supplied to overcome the energy barrier of the nucleation [5, 6]. During devitrification transformation a very high nucleation frequency occurs with limited grain growth resulting multiple nano-crystalline phases [7, 8].

Microstructural evaluation in these kinds of steels showed that nanosize phases i.e., Fe5C12Mo10, Fe3B, Fe3C and α-Fe were nucleated in the amorphous structures after annealing process [9]. It was also proof that by nucleating of mentioned nanosize phases not only hardness and wear resistance but also abrasion and fretting re-sistance of the alloys were promoted erratically [10, 13].

Although several researches are devoted to characterization of extreme hardness steels achieving from bulk amorphous steels, there are only a few attempts toward the detailed studies of the crystallization kinetics of alloys. In this paper, we launched the research toward expanding the previous works by D. J. Branagan [14] and his co-workers to evaluate the key factors in developing nanostructured ultra-high hardness steels categorized at metal matrix nanocomposites. To facilitate this idea, chemical composition of the alloys was designed on the class of materials called hardmetals. In fact, in this research work effects of alloying process via transient elements (Nb, Mo, Cr) on hardening of Fe-based nanostructured alloys have been studied. In other words, achieving high hardness, more than 18 GPa, in a nanostructured Fe-based composite alloy was evaluated in the research.

2. METHOD AND MATERIALS

In this research study, thin ribbons prepared from as cast cube ingots were used for kinetic and hardness investigations. In fact, multi-component Fe-based alloy ingots were prepared in an arc
furnace with nominal compositions of Fe<sub>50</sub>Cr<sub>10</sub>Mo<sub>25</sub>B<sub>15</sub>Cr<sub>2</sub>Nb<sub>5</sub>. Pure iron (99.7 mass %), chromium (99.9 mass %), niobium (99.9 mass %), molybdenum (99.9 mass %), and crystalline B (99.5 mass %) were used in an argon atmosphere to produce ingots. To achieve fully amorphous structures, rapidly solidified thin ribbons with a thickness of about 60 μm were prepared by melt-spinning technique (wheel speed: 32 m/s). Then, amorphous ribbons were annealed under vacuum (10<sup>-3</sup> torr) in a furnace above the crystallization temperatures as the annealing process to obtain nanostructured structures.

Philips XRD device with CuKα radiation (voltage: 40 kV, amperage: 40 kA), equipped with Xpert software was used to identify crystallized phases. Composition of the ribbons was verified by using energy-dispersive X-ray spectroscopy. Moreover, microstructural evaluations were accomplished by a 200 kV JEOL transmission electron microscope equipped with an energy dispersive X-ray spectrometer (INCA PentaFETX3-Oxford instruments). Before applying microstructural determination, samples were prepared in foil shape with 3 mm diameter. These discs were electro polished in twin jet electro polisher with a sol-vent composed of 25% nitric acid and 75% meth-anol.

Vickers microhardness measurements were done with a 30 g load on the cross section of the samples using a Reichert Jung Micro Durmatt 4000 E system. For each sample, 7 to 10 hardness indentations were made and then the average of the measurements was reported. The average standard deviation for the hardness measurements of the ribbons was 0.385 GPa.

3. RESULTS AND DISCUSSION

3.1. TEM Observations

In figure 1, XRD pattern of the alloy (after melt-spinning process) is shown. As can be seen clearly, there are no significant crystalline sharp peaks in the chart showing an amorphous structure (formed in the samples after melt-spinning process).

In figure 2 result of the X-ray diffraction test is shown determining crystallization of α-Fe and other crystalline phases in the structure of the alloy after annealing process. It is clear that after annealing process structure of the alloy was consist of crystalline phases such as α-Fe, Fe<sub>3</sub>C, Fe<sub>23</sub>B<sub>6</sub>, Fe<sub>2</sub>B and Fe<sub>50</sub>Cr<sub>12</sub>Mo<sub>10</sub>. In fact multi-phase structure was given after the heat treatment.

In figures 3 to 6 microstructures of the alloy in amorphous and also crystalline states are shown.

Significantly no crystalline phases are detected in the amorphous state (figure 3). It is essential to mention that one of the most striking features of the Fe-based nanostructured alloys produced by heat-treating of a preliminary amorphous state is the distinct morphologies of the crystalline phases (i.e. α-Fe, Fe<sub>3</sub>C, Fe<sub>23</sub>B<sub>6</sub>, Fe<sub>2</sub>B and Fe<sub>50</sub>Cr<sub>12</sub>Mo<sub>10</sub>).

![XRD pattern](image-url)  
**Fig.1.** XRD pattern of Fe<sub>50</sub>Cr<sub>10</sub>Mo<sub>25</sub>B<sub>15</sub>Cr<sub>2</sub>Nb<sub>5</sub> in amorphous state.
In the figures, microstructures of the alloy after heat-treating above the second crystalline temperature are shown. One can be seen that α-Fe crystals were formed in the structures of the alloys in mottled morphology. As mentioned earlier, unique features of crystalline phases can be effectively used toward identification of these nano-size phases. The nano scale phases crystallized in the structure of the alloy in special morphologies mentioned as following:

i. α-Fe: mottled structure
ii. Fe₃B: multi-twinned structure
iii. Fe₅C: pentangle structure
iv. Fe₁₀Cr₁₂Mo₁₆: layer (perlitic) structure

3. 2. Hardness Evaluation

Hardness measurements were taken on the cross section of both amorphous and heat treated ribbons; results are given in table 1. In figure 7, an example of the hardness identification on the cross section of a heat treated is shown. The as-cast ribbon exhibited high level of hardness of 13 GPa which become harder after heat treatment. Despite the similar alloys investigated by D. J. Branagan [14] and some other works [15, 16], in this new alloy amorphous samples show higher
level of hardness that those of previous works. On the other hand, hardness significantly increases in the samples after heat treating but in the hard metals evaluated since now, similar observations have not been reported.

With respect to the crystallization of the nanoscale iron based phases, hardness mechanism in these nano-composite steel alloys are significantly different with that found in conventional steel alloys. In fact in the amorphous state, the key factor toward the extreme hardness is related to solid solution strengthening through non-equilibrium solid state. Due to the high amount of the alloying elements (i.e. Cr, Mo, Nb, and B) supper saturated solid was formed during rapid quenching. In other words, super saturations of the transition metallic elements near and above their equilibrium solubility limits are the first key factor in attending extreme hardness in amorphous state.

The second key factor toward obtaining high level of hardness is reduction of structure from amorphous solid to nano-crystalline solid. Indeed, hardness in the alloy was developed upon 18.6 GPa while crystal size decreased to about 50nm by heat treating over the crystallization temperature. By considering the data given in table 1, it can be inferred that forming supper saturated solid (amorphous state) and then converting this solid to the crystalline structure consisting of the nano-size phases are the most effective mechanisms toward hardening of the alloy.

4. CONCLUSIONS

The hardness levels obtained by devitrification process in the alloy (18.6 GPa) are more than those of the similar hardmetals. Nb alloying is hypothesis as the key factor toward increasing hardness in both amorphous (by supersaturation of alloying elements e.g. Nb) and crystalline state (by reduction in the structure to the nanometer phases).

REFERENCES

Table 1. Results of hardness tests in the alloy

<table>
<thead>
<tr>
<th>alloy</th>
<th>Hardness in amorphous state (GPa)</th>
<th>Hardness in crystalline state (GPa)</th>
<th>Average grain size (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe&lt;sub&gt;0.6&lt;/sub&gt;Cr&lt;sub&gt;1.4&lt;/sub&gt;Mo&lt;sub&gt;0.5&lt;/sub&gt;B&lt;sub&gt;1.2&lt;/sub&gt;C&lt;sub&gt;4&lt;/sub&gt;Nb&lt;sub&gt;0.5&lt;/sub&gt;</td>
<td>13</td>
<td>18.6</td>
<td>50</td>
</tr>
<tr>
<td>[Fe&lt;sub&gt;0.8&lt;/sub&gt;Cr&lt;sub&gt;0.2&lt;/sub&gt;]&lt;sub&gt;7.6&lt;/sub&gt;W&lt;sub&gt;1.2&lt;/sub&gt;C&lt;sub&gt;1.8&lt;/sub&gt;</td>
<td>11</td>
<td>16.2</td>
<td>150</td>
</tr>
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