AISI S1 tool steel after deep cryogenic treatment: Tensile properties and microstructure

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Abstract: Successful employment of advanced tool steel in engineering applications is related to its ability in terms of meeting service life requirements and fabrication with proper dimensions. Deep cryogenic treatment may be used to produce advanced tool steel by simultaneously increasing toughness, strength, and hardness. Twelve sets of specimens were tested, 9 of which were deep cryogenic treated and then tempered. Tensile properties, hardness, X-ray diffraction, and scanning transmission electron microscopy were applied for macroscopic and microscopic investigations. The best results of simultaneous improvement in tensile toughness, hardness, and strength were obtained for 36 h soaking and 1 h tempering times.

Keywords: Hardness, STEM, Strength, Tensile Toughness, XRD

1. Introduction

Sustainable development is an important challenge in human life. It refers to changing the process of proper application of resources, conducting investments, and orienting technology development in a way that could be consistent with present and future needs. Proper use of raw materials facilitates sustainable development [1-3]. Advanced tool steel properties include high strength, wear resistance and hardness in addition to suitable toughness for enduring mechanical impact loadings. Researchers [4-9] demonstrated that sub zero treatments improved mechanical properties of several steels. For example, for AISI M2 high speed tool steel hardness and wear resistance improved by using DCT compared with the conventional treatment [6]. Therefore, DCT used in this study for producing advanced AISI S1 tool steel.

2. Experimental methods

Table 1 demonstrates the chemical analysis of AISI S1 used in this study. The AISI S1 is mostly used for manufacturing cutting blades for thick sheets and also for punching tools.

<table>
<thead>
<tr>
<th>Element</th>
<th>(W%)</th>
<th>Element</th>
<th>(W%)</th>
<th>Element</th>
<th>(W%)</th>
<th>Element</th>
<th>(W%)</th>
<th>Element</th>
<th>(W%)</th>
<th>Element</th>
<th>(W%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>0.4800</td>
<td>Ni</td>
<td>0.1280</td>
<td>S</td>
<td>0.0250</td>
<td>W</td>
<td>1.5700</td>
<td>Mo</td>
<td>0.0281</td>
<td>Fe</td>
<td>Rest</td>
</tr>
<tr>
<td>Si</td>
<td>0.9950</td>
<td>V</td>
<td>0.0148</td>
<td>Cr</td>
<td>1.1200</td>
<td>Mn</td>
<td>0.3360</td>
<td>P</td>
<td>0.0567</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Many studies have been published on advanced tool steels; however, each one has focused on one or few sets of specimens rather than collecting data from many specimens. Consequently, it is helpful to collect the published results in order to have thorough representations of the measured effects. Thus, 12 sets of AISI S1 specimens were tested. One set consisted of the specimens which underwent the conventional hardening procedure [10]. The remaining sets of specimens underwent full treatment cycles. The temperature-time history for the specimens is given in Fig. 1(a) and flowchart of the experimental procedure is depicted in Fig. 1(b). The DCT work followed a procedure shown in Fig. 1(b) and used a programmable cryogenic processor which was designed and developed [11]. Considering the
required tempering temperature of 200ºC, the cryogenic processor had some added facility for the in-situ tempering of the DCT specimens for the required time durations. The time from the end of hardening to the beginning of DCT was ≈17 min. However, the DCT specimens were tempered in-situ, which led to the minimum time gap between the steps as ≈2 min. The specimens were assigned codes for easy identification, which is described in Tables 2 and 3.

The specimens had to be machined to the required sizes prior to their treatment. The uniaxial force-displacement tensile test (F-∆L) was carried out on a standard specimen with 5 mm diameter and 25 mm parallel length at room temperature, according to BS EN 10002-1 [12]. The strain rate was 0.00166 S-1, each test was repeated for three times, and the average value was obtained. For example, it depicted for specimens of 361 and 003 in Fig. 2.

A fast and reliable tool for phase and orientation map acquisition by an STEM was obtained in [13]. STEM specimens were cut out as 3 mm discs in the wire cut machine, thinned to ~100nm by hand polishing, then dimpled and argon ion milled using a Gatan PIPS mill and cold stage. STEM machine used at LeRoy Eyring Center for Solid State Science, John Cowley Center for HREM, Arizona State University, USA. Theoretical patterns calculated for the phases listed in the ASTM E975-13 [14].
3. Results and Discussion

3.1. Tensile Evaluations

As shown in Figs. 3 and 4, for different specimens, yield strength and UTS showed no relative changes. However, in comparison with the standard specimen, UTS and hardness of the DCT specimens had 32-36% and 9-12% increase, respectively.

Fig. 5 shows that, with increasing the tempering time, the soaking time required for obtaining the maximum hardness was reduced. For example, for 1 h of tempering time, 48 h of soaking time at -196ºC was needed to achieve maximum hardness. For 2 h of tempering time, the corresponding soaking time at -196ºC was approximately 24 h and further increase in soaking time was not necessary. In other words, with increase in tempering time, shorter soaking time was required for obtaining maximum hardness. In addition, Fig. 5 demonstrates that, with increase in soaking time, the tempering time required for obtaining maximum hardness was reduced. For example, for 24 h of soaking time at -196ºC, 3 h of tempering time was needed to achieve maximum hardness. For 36 h of soaking time at -196ºC, the corresponding tempering time was 2 h. In the case of 48 h of soaking time, the required tempering time was 1 h. In other words, with increase in soaking time, shorter tempering time was required for achieving maximum hardness.

The above findings could be explained by the results of microstructure section. Volume fraction (Vf) of SC increased with soaking or tempering times from 2.18 V% to 12.87 V% (Fig. 10), which led to maximum hardness.

As demonstrated in Fig. 6, among all the specimens, elongation of specimen 242 was the minimum, while specimens 361 and 482 had maximum elongation. Therefore, according to Figs. 4 and 6 and considering the following relation in [15] (Equation 1), specimens 361 and 482 presented 12-35% improvement in tensile toughness as demonstrated in Fig. 7; where UTS is ultimate tensile strength and εf is fracture strain.

$$\frac{2}{3} \times \text{UTS} \times \varepsilon_f \approx U_T$$ or tensile toughness \hspace{1cm} (1)
As demonstrated in Fig. 7, in this research, specimens 361 and 482 had higher tensile toughness than conventionally treated ones, which showed that increasing DCT duration needed increasing tempering time to obtain high tensile toughness. For example, for 36 h of soaking time at -196°C, 1 h of tempering time was needed to achieve maximum tensile toughness. For 48 h of soaking time at -196°C, the corresponding tempering time was 2 h. In other words, with increase in the tempering time, longer soaking time was required for obtaining maximum tensile toughness, because tempering after DCT caused more reduction of residual stress than tempering after conventional treatment [16]. In other words, increasing DCT duration produced more dislocations [17, 18], which needed increasing tempering time (at constant tempering temperature) to obtain high tensile toughness.

Therefore, at -196°C, the retained austenite was completely transformed into martensite, which confirmed the results from Cu Ka radiation diffraction (XRD) phase analysis PHILIPS at 40KV. As demonstrated in Fig. 8 for specimens 002 and 483, the retained austenite was not observed.

Fig. 9(a) shows that there was no dislocation around the carbide particles. Also, EDS analysis showed this particle was iron carbide. Iron carbide diffraction patterns were sensed near the retained austenite grains; but, there was no compositional signal. Fig. 9(b) demonstrates spherical shape carbides with sub-micron sizes. Also, SCs were dispersed, as can be seen in this low magnification micrograph.

Fig. 9(c) shows STEM map of M2C6, M23C6, Fe3C, M6C, and Cr7C3 carbides and retained austenite in the ferrite matrix, which was taken from the same area of about 80000 nm². This method detected new carbides (Fe3C, M2C and Cr7C3) which could not be detected by SEM in the previous research [17]. According to the second mechanism of precipitation of carbides (SCs or TCs) [17], during DCT, there would be appropriate places around PCs for the precipitation of carbides (SCs or TCs). At low tempering temperature, diffusion was more difficult. So, the question is: how does it work? First, diffusion coefficient of carbon is higher than alloying element ones (such as W and Cr). Second, in grain boundaries, diffusion coefficient is higher than that inside the grains. All of these points would lead to the facilitated precipitation of carbides (SCs or TCs) around the PCs in grain boundaries.
(a) High Resolution Brightfield STEM

- Retained Austenite in this region.
- Dark Squiggly Lines are Dislocations.

(b) HAADF (high angle annular dark field) STEM

- Brightfield STEM
As depicted in Fig. 10, volume fraction of the PC was approximately constant (≈0.4±0.2 V%) in different specimens, while Fig. 10 demonstrates that volume fraction of the SC increased with soaking or tempering times from 2.18 V% to 12.87 V%. As represented in Fig. 10, an increase in soaking time resulted in the precipitation of more SC (50% on average) due to dislocation density around the PCs in phase matrix, which increased with an increase in the soaking time and facilitated SC precipitation. It can be observed in Fig. 10 that an increase in tempering time resulted in the slight precipitation of SC (on average 50%). On the other hand, volume fraction of SC in specimens 002 and 003 was somehow fixed, since, without carrying out DCT, the conditions for the formation of SC were not facilitated.

According to Fig. 11, population density (PD) of PC was constant for all the specimens. Also, Fig. 12 shows that average size of the PC remained almost constant with increase in soaking or tempering times (especially, for long soaking times of 48 h, for example), which was the specifications of PC depending on the austenitization treatment parameters that were kept constant in this research.

According to Fig. 11, in the specimens with similar soaking times, PD of SC was reduced with increase in the tempering time, which was due to the fact that, for the same soaking time, density of the dislocation which was made during the process was almost the same. Therefore, with an increase in tempering time, growth of SCs would take place owing to their joining because of relatively shorter distance.

As shown in Fig. 12, with increase in the tempering time, size of SC had a slight increase (up to 13.6%). Similarly, according to Fig. 12, with an increase in the soaking time, size of SC also increased (up to 100%) due to the increase in dislocation density. Additionally, lattice distortion in the regions surrounding SC was relatively higher due to the relatively less reduction in the alloy elements and carbon content owing to being utilized in the precipitation of relatively fine SC.

Comparative assessment of the characteristics of SCs of
specimen 361 and those of specimens 002 and 003 could help infer that volume fraction of SCs increased by 125% (Fig. 10) and PD of SCs increased by 350% (Fig. 11) which was associated with reduction in their mean diameter by 30% (Fig. 12). These results unambiguously indicated that DCT refined SCs.

4. Conclusions

Different soaking and tempering times were considered to study the effect of various treatment cycles on tensile properties and microstructure of advanced AISI S1 specimens. Kinetically, in the present research, the most appropriate times were 36 h and 48 h of deep cryogenic and 1 h and 2 h of tempering treatments, respectively, because 12-35% improvement in tensile toughness, 9-12% improvement in hardness, and 32-36% improvement in strength were obtained at these times. For advanced AISI S1 steel, it was observed that maximum tensile toughness could be obtained by simultaneously increasing soaking and tempering times. But, if reduction in production time and cost was of primary concern, lower soaking and tempering times would seem to be a more feasible option. In this case, 36 h of soaking time and 1 h of tempering time would be a more feasible option.

Nomenclature

DCT Deep Cryogenic Treatment
PD Population Density
PC Carbide precipitates as the consequence of the solidification
Secondary Carbide
SC Carbide precipitates as the consequence of the first tempering
SEM Scanning Electron Microscopy
STEM Scanning Transmission Electron Microscopy
VF Volume Fraction
XRD X-ray diffraction

References


Figure 12. Average size of the PCs and SCs