International Journal of Minerals, Metallurgy and Materials Volume 22, Number 3, March 2015, Page 272 DOI: 10.1007/s12613-015-1071-5

# Microstructural evolution of a heat-treated H23 tool steel

*Meilinda Nurbanasari*<sup>1)</sup>, *Panos Tsakiropoulos*<sup>2)</sup>, and Eric J. Palmiere<sup>2)</sup>

 Department of Mechanical Engineering, Institut Teknologi Nasional, Bandung, JI. PHH. Mustapha 23, Bandung, 40124, West Java, Indonesia
 Department of Materials Science and Engineering, The University of Sheffield, Sir Robert Hadfield Building, Sheffield S1 3JD, United Kingdom (Received: 22 May 2014; revised: 3 October 2014; accepted: 8 October 2014)

Abstract: The microstructure and the stability of carbides after heat treatments in an H23 tool steel were investigated. The heat treatments consisted of austenization at two different austenizing temperatures ( $1100^{\circ}C$  and  $1250^{\circ}C$ ), followed by water gene ing and double-aging at 650°C, 750°C, and 800°C with air cooling between the first and second aging treatments. Martensite did not form in the as-quenched microstructures, which consisted of a ferrite matrix, M<sub>6</sub>C, M<sub>7</sub>C<sub>3</sub>, and MC carbides. The double-aged microstructures consisted of a ferrite matrix and MC, M<sub>6</sub>C, M<sub>7</sub>C<sub>3</sub>, and M<sub>23</sub>C<sub>6</sub> carbides. Secondary hardening as a consequence of secondary productions. The highest double-aged hardness of the H23 tool steel was 448 HV after austenization at 1250°C and double-aging at 650°C, which suggested that this tool steel should be used at temperatures below 650°C.

Keywords: tool steel; microstructural evolution; heat treatment; carbides; stability; had

# 1. Introduction

The high concentration of carbide-forming ts in tool steels encourages the formation of a carbine network in the as-cast microstructure, which can d crease the steels' toughness. The type of carbide, its portuge fraction, morphology, size, and distribution can be modified through heat-treatment processes to achieve the required mechanical properties. H23 tool steel with is one of the tungsten hot-work tool-steel roups, cortains high concentrations of chromium and tungs and is used for applications at elevated working temperatures (about 600°C) such as medium runs on brass die-casting dies. The H23 tool steel also has good hot strength and resistance to heat checking [1]. Previous studies [2-6] on the microstructure evolution and properties of hot-work tool steels after heat treatment showed a strong correlation between carbide precipitation and mechanical properties. Because the carbide affects the mechanical properties of the tool steel, the microstructure and types of carbide become very important. The carbide precipitation process during aging can be complicated when steels contain more than one carbide-forming element. Thomson and Bhadsehia [7] suggested that the sequence of carbide formation in high-chromium steels is matrix  $\rightarrow M_3C$  $\rightarrow M_7C_3 \rightarrow M_{23}C_6$ . Dudova and Kaibyshev [8] stated the carbide sequence in a 10% chromium steel is martensite  $\rightarrow M_3C \rightarrow M_2C \rightarrow M_{23}C_6$  and M<sub>6</sub>C. Asadabad *et al.* [9] studied the aging of a 4.5%Cr–2%W–0.25%V (wt%) steel and observed that the M<sub>3</sub>C and M<sub>2</sub>C carbides formed in the first stages of aging and subsequently transformed into the M<sub>7</sub>C<sub>3</sub> and M<sub>23</sub>C<sub>6</sub> carbides.

Although numerous previous studies on tool steels have been reported [2–6], the microstructure changes, carbide precipitation, and mechanical properties of the heat-treated H23 tool steel have not been established. The difference in chemical composition and heat treatment of tool steel will also result in different microstructures and carbides that affect the mechanical properties of tool steels. Hence, in this study, we contribute to this growing area of research by exploring the microstructure and the stability of the carbide in H23 tool steel after heat treatment. We then compare the experimental results with those predicted using thermodynamic calculations.



Corresponding author: Meilinda Nurbanasari E-mail: meilinda@itenas.ac.id, nurbanasarimeilinda@yahoo.com © University of Science and Technology Beijing and Springer-Verlag Berlin Heidelberg 2015

# 2. Experimental

A vacuum induction melting furnace was used to produce a square ingot of an H23 tool steel with nominal composition: 0.3%C-12.3%W-12.3%Cr-1.2%V-0.4%Mn-0.4%Ni-Fe balance (wt%) [10]. The 1-cm<sup>3</sup> samples for heat treatment were cut from the middle top of the ingots using an electric discharge machine. The samples were heated at 1100°C and 1250°C under an argon atmosphere with a 1-h holding time at each temperature and were subsequently water quenched. The water quenching was used to avoid grain coarsening and to achieve the optimum hardness. Double-aging was then conducted at three different aging temperatures of 650°C, 750°C, and 800°C, with a 1-h holding time at each temperature. Air cooling was performed between the first and second aging treatments. Groesbeck's etchant (100 mL of H<sub>2</sub>O, 10 g of NaOH, and 10 g of KMnO<sub>4</sub>) was used to reveal the carbides. The etching time was approximately 5-10 s. The optical microstructure observations were performed using an MET Polyvar microscope. The scanning electron microscopy (SEM) investigations were conducted on Inspect F (20 kV) and JEOL 6400 microscopes (20 kV), the latter of which was equipped with an energy-dispersive X-ray spectroscopy (EDS) unit; the INCA software was used for quantitative chemical analysis. All carbides larg than 0.1  $\mu$ m as well as their volume fraction (V<sub>c</sub>) based in SEM images were calculated using the ImageI\_\_\_\_ Thin foils for transmission electron microstopy (JEM) studies were prepared using electrolytic twinget polishing with a solution of 5vol% perchloric cia, 35vol% butoxyethanol, and 60vol% methanol. The TENI studies were conducted on a JEOL 2010F microscology (200 kV) equipped with an EDS unit operated using the Oxford ISIS software. A CV Instrument Vickers Largers' tester was used to measure the hardness with a load of 10 kg. Phase and carbide identification were performed on a Siemens D5000 X-ray diffractometer equipped with a Co radiation source. The polished samples were scanned over the  $2\theta$  range of  $30^{\circ}$  to 130° at a 0.02° step size and a 1° per minute counting time. The STOE WinXPow software program and the ICDD (International Centre for Diffraction Data) files were used for peak identification. The calculated phase diagram of the H23 tool steel was obtained using the ThermoCalc software (TCEF6 database) [10].

# 3. Results and discussion

### 3.1. As-cast microstructure

Nurbanasari et al. [10] reported that the bulk hardness of

cast H23 tool steel was  $355 \pm 4$  HV and that the as-cast microstructure consisted of a ferrite matrix and carbides, as shown in Fig. 1. The carbides in the as-cast H23 tool steel were coarse tungsten-rich M<sub>6</sub>C carbide, fine vanadium-rich MC carbide, and fine chromium-rich M<sub>23</sub>C<sub>6</sub> carbide.

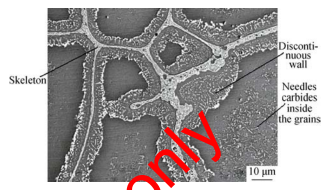


Fig. 1. Secondary-electron in age of the as-cast H23 tool steel [10].

# 3.2. Effects of stenizing temperature

of austenizing temperature on microstructure 3.2.1. ffee Fig. 2 shows the as-quenched microstructures of the H23 of teel prepared at two different austenizing temperatures. Ferrite and carbides coexisted, and no martensite was observed in the samples prepared at either austenizing temperatures because the Ms temperature of this tool steel is approximately -46°C [1]. Compared with the as-cast microstructure (Fig. 1), dissolution and separation of the carbides were observed in both austenized steels. In the sample austenized at the lower temperature, the carbides were still interconnected in the thick layer (skeleton) and carbides in the discontinuous wall dissolved. In the sample austenized at the higher temperature, the carbides in the skeleton became separated and coarser and the carbides in the discontinuous wall dissolved into the matrix to a greater extent. The microstructure of the as-quenched H23 tool steel austenized at the higher temperature exhibited a lower volume fraction of carbides because more carbide dissolved into the matrix. In the steel austenized at the higher temperature, a large volume fraction of undissolved M6C carbides remained. The microstructures of the as-quenched H23 tool steel exhibited no indication of the inside-the-grain carbide needles observed in the as-cast steel (Fig. 1). These small needle carbides dissolved more easily into the matrix than the larger carbides because of their greater surface-to-volume ratio; i.e., smaller carbides were less stable than larger carbides of the same carbide type.

Figs. 3 and 4 show SEM images and EDS spectra of the matrix and carbides of the as-quenched H23 tool steel. Some

carbides exhibited two different contrasts under back scattered electron imaging conditions, which indicated that two types of coarse carbides formed in the as-quenched H23 tool steel. These carbides were identified by SEM–EDS as the M<sub>6</sub>C (bright) and M<sub>7</sub>C<sub>3</sub> carbides (grey), and this identification was supported by the XRD results (see Fig. 5), which confirmed the presence of  $\alpha$ -Fe–Cr (the super-saturated solid solution), austenite, and the M<sub>6</sub>C and M<sub>7</sub>C<sub>3</sub> carbides. The M<sub>6</sub>C carbide was Fe<sub>3</sub>W<sub>3</sub>C, and the M<sub>7</sub>C<sub>3</sub> carbide was (Cr,Fe)<sub>7</sub>C<sub>3</sub>. Additional and more intense peaks associated with the Fe<sub>3</sub>W<sub>3</sub>C and (Cr,Fe)<sub>7</sub>C<sub>3</sub> carbides were observed in the EDS spectra of the sample at austenized at 1100°C than in the spectrum of the sample austenized at 1250°C, which indicated that the volume fraction of these carbides was greater at the lower austenizing temperature. The following discussion shows that the dissolution rates of  $M_6C$  and  $M_7C_3$ carbides differ and that a higher austenizing temperature results in the dissolution of more carbides into the matrix. No peaks associated with the  $M_{23}C_6$  carbides were observed in the EDs spectra. Typical EDS spectra of the  $M_6C$  carbides (Figs. 3 and 4) showed intense tungsten and iron peaks. The typical EDS spectra of the  $M_7C_3$  carbides exhibited an iron peak more intense than the chromium peak or vice versa [11–12]. Both types of carbides exhibited rod-like, irregular,

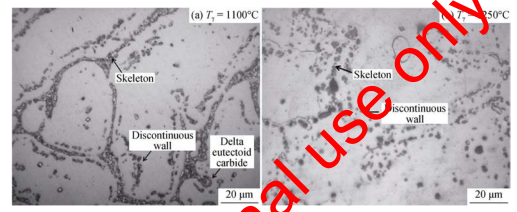


Fig. 2. Optical images of the microstructure of the F25 tool steel after austenization and water quenching.

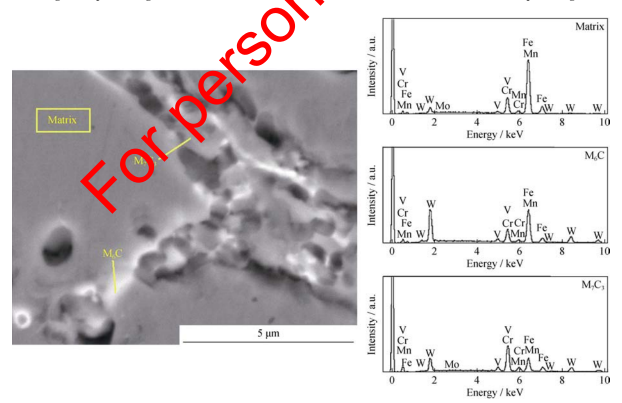


Fig. 3. Backscatter electron image of the H23 tool steel water quenched from  $T_{\gamma} = 1100^{\circ}$ C and EDS spectra of the matrix and carbides.

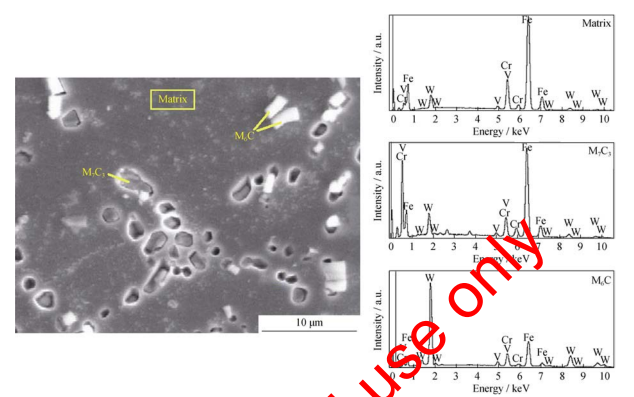


Fig. 4. Backscatter electron image of the H23 tool steel water quenched from  $T_{\gamma} = 1250^{\circ}$ C and EDS spectra of the matrix and carbides.

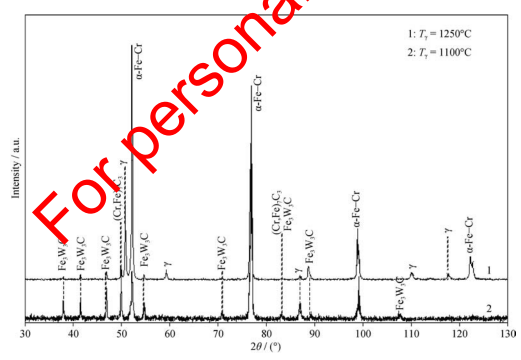


Fig. 5. X-ray diffraction pattern of the as-quenched H23 tool steel. The ICDD cards used were those of  $\alpha$ -Fe–Cr (34–396), austenite (52–512), Fe<sub>3</sub>W<sub>3</sub>C (41–1351), and (Cr,Fe)<sub>7</sub>C<sub>3</sub> (5–720).

and spherical morphologies. The presence of the  $M_{23}C_6$  carbide in the as-quenched H23 tool steel austenized at both temperatures was not observed by either SEM or TEM. The

 $M_{23}C_6$  carbides easily coalesce and coarsen during austenization and could dissolve completely at about 1095°C [13–14]. Thus, the primary  $M_{23}C_6$  carbides (i.e., the carbides that formed upon solidification) dissolved gradually during austenization, thereby enriching the ferrite matrix with chromium and carbon and resulting in nucleation of the  $M_7C_3$  carbide [15–16].

The quantitative data of the matrix and carbides measured using SEM–EDS are summarized in Table 1. The compositions of the matrices were in good agreement with those calculated using ThermoCalc-TCFE6 (79.9%Fe– 13.1%Cr–1.1%V–5.9%W (wt%) at  $T_{\gamma} = 1100^{\circ}$ C and 76.9%Fe–12.6%Cr–1.1%V–9.4%W (wt%) at  $T_{\gamma} = 1250^{\circ}$ C). Table 1 also shows that the M<sub>7</sub>C<sub>3</sub> carbide was rich in iron and chromium and that its chromium content decreased with increasing austenizing temperature. The diffusivity of alloying elements increases with increasing temperature and the diffusivity of chromium is higher than that of tungsten. Because the  $M_7C_3$  carbide was rich in chromium, its dissolution was affected by its chromium content that reached its maximum value (and thus maximized the stability of the  $M_7C_3$  carbide) at the austenizing temperature of 1100°C. The austenizing temperature at which the  $M_7C_3$  carbide dissolves is higher than that at which the  $M_2C_3$  carbide dissolves [17]. The dissolution rate of the  $M_7C_3$  carbides was faster than that of the  $M_6C$  carbides, and the  $M_7C_3$  carbides to dissolve at the higher austenizing temperature, where their chromium content decreased. This result contrasted that for the  $M_6C$  carbides, where the tungsten content increased with increasing austenizing temperature.

	Table	1. SEM–EDS data o	of phases in the as-q	uenched H23 tool	steel	wt%
T/9C	Phase		C			
$T_{\gamma} / ^{\circ}\mathrm{C}$		Fe	Cr	V	W	Comment
	Matrix	$78.7\pm0.6$	$13.8\pm0.6$	11+01	$6.5\pm0.2$	Ferrite
1100	Carbide	$51.9\pm5.7$	$13.0 \pm 1.1$	2.5 ± 1 2	$32.4 \pm 3.4$	M <sub>6</sub> C
	Carbide	$50.0\pm1.2$	27.7 ± 0.3	$-3 \pm 0.2$	$19.1\pm0.3$	$M_7C_3$
	Matrix	$75.0\pm0.9$	$14.0 \pm 0.2$	$1.2 \pm 0.0$	$9.8\pm0.6$	Ferrite
1250	Carbide	$31.1 \pm 6.6$	9.1 - 0.	$2.6 \pm 0.2$	$56.9 \pm 3.3$	M <sub>6</sub> C
	Carbide	$76.1\pm1.9$	112±2	$1.1\pm0.1$	$11.6\pm0.9$	$M_7C_3$

Note: \* The quantitative data should be considered with caution given the size of the carbides.

Using SEM–EDS with accelerating voltage 20 kV requires a particle size greater than 1  $\mu$ m for the inalyzed carbide to obtain proper quantification withou overlap from another constituent in the interaction volume. The quantitative data in Table 1 were taken from cardides with sizes in the range 0.3–1.5  $\mu$ m. Thus, interference between the interaction volume of the matrix and the smaller carbides was unavoidable. In addition, on the basis of data in Table 1, a minor difference existed in the compositions of the alloying elements between the M<sub>7</sub>C<sub>3</sub> carbides and the matrix at 1250°C because of the interference of the relatively large interaction volume with the matrix.

TEM studies were conducted on the sample austenized at the higher temperature to compliment the as-quenched microstructure characterization (see Fig. 6). The diffraction patterns of the austenite, ferrite, and MC carbide in Fig. 6 confirmed that these phases were present in the as-quenched microstructure. Because this tool steel contained high concentrations of ferrite-stabilizing elements (chromium and tungsten) and because its Ms temperature is far below room temperature [1], ferrite unsurprisingly existed in the as-quenched microstructure. The retained austenite observed in bright-field TEM images was a film type, with a width 10 to 60 nm and located at the ferrite lath boundaries. The formation of retained austenite was due to the dissolution of the carbides, which resulted in locally high carbon content in the matrix; water quenching from the higher austenizing temperature did not provide sufficient time for the austenite to completely transform to ferrite. The size of the MC carbides was about 40 to 50 nm. The MC carbide and the ferrite matrix exhibited the Baker–Nutting relationship [18]. 3.2.2. Effect of austenizing temperature on hardness

The hardness of the as-quenched tool steel was  $279 \pm 1$  HV and  $302 \pm 7$  HV after being quenched from austenizing temperatures of 1100°C and 1250°C, respectively. The as-quenched hardness of the tool steel was lower than the hardness of the cast steel because the austenite did not transform to martensite. The higher hardness of the as-quenched tool steel austenized at the higher temperature was attributed to the dissolution of carbides into the matrix, which enriched the matrix with alloying elements. The high tungsten and chromium contents in the H23 tool steel stabilized the presence of delta ferrite at the austenizing temperature. This ferrite was stable to room temperature and

did not contribute to hardening. Thus, the lower hardness with decreasing austenizing temperature was not only due to the lower dissolution of carbides, but was also due to the higher volume fraction of ferrite and the non-transformation of austenite to martensite because of the low Ms temperature.

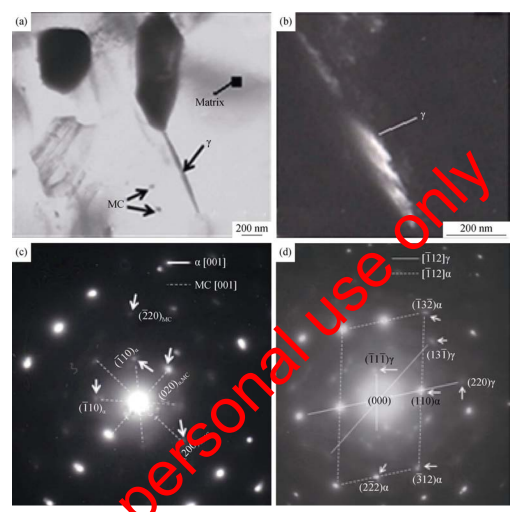


Fig. 6. Top row: bright-field TFM image showing MC carbides, ferrite (matrix), and retained austenite in the H23 tool steel austenized at 1250°C and water querched (a) and dark-field TEM image showing retained austenite (b). Bottom row: selected-area diffraction patterns (SADBs) of the MC carbide in the ferrite matrix (c) and the retained austenite and ferrite (d).

Table 2 shows the time austenizing temperature strongly influenced the volume naction of carbides, which decreased with increasing austenizing temperature, and subsequently enriched the matrix with alloying elements via the dissolution of carbides, whose extent of dissolution increased in the

Table 2.Volume fraction of carbides and their average size inthe H23 tool steel water quenched from different austenizingtemperatures

A a post	andition	As quenched condition					
As cast condition		$T_{\gamma} = 1$	250°C	$T_{\gamma} = 1100^{\circ} \text{C}$			
V <sub>c</sub> / %	Mean	V <sub>c</sub> / %	Mean	V <sub>c</sub> / %	Mean		
	size / µm	. (, , , ,	size / µm	. (, , , ,	size / µm		
$21.5\pm0.4$	$0.4\pm0.1$	$13.9\pm1.0$	$0.5\pm0.1$	$15.3\pm1.9$	$0.4 \pm 0.2$		

sequence MC,  $M_6C$ ,  $M_7C_3$ ,  $M_{23}C_6$  [19]. The correlation between the mean size of carbides and the austenizing temperature was not conclusive because the mean size of carbides was measured for individual carbides and the carbide networks still existed at the lower austenizing temperature.

# 3.3. Effect of double-aging treatment

#### 3.3.1. Effect of double-aging treatment on microstructure

The microstructure evolution of tool steels is distinctly affected by the different aging conditions. Fig. 7 shows the microstructure of the H23 tool steel after treatment at different austenizing and double-aging temperatures. The double-aged microstructures austenized at different temperatures consisted of ferrite and carbides. In general, the carbides were more separated in their prior network after austenization at the higher temperature and were still interconnected in their prior network after austenization at the lower temperature; in addition, the undissolved delta ferrite was also observed in the sample austenized at the lower temperature. Fig. 7 also shows that, in the samples austenized at the higher temperature, fine secondary carbides were observed after the double-aging treatment at 650°C; in contrast, in the sample austenized at the lower temperature, fine secondary carbides were observed under double-ageing at 750°C. The volume fraction of secondary carbides was higher in the samples austenized at the higher temperature than in the samples austenized at the lower temperature when the same double-aging temperature was used. This greater volume fraction was due to the increased dissolution of carbides at the higher austenizing temperature causing more carbon to be dissolved in the matrix, which provided the driving force for carbide precipitation during the double-aging process [6].

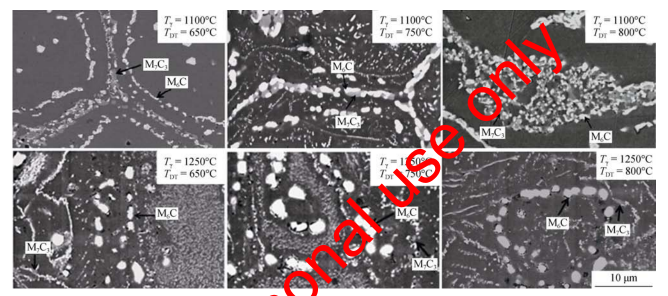


Fig. 7. Backscatter electron images of the H2. 460 steel austenized and double-aged at various temperatures.

The presence of retained austenite after wave quenching also affected the secondary carbide precivitation process during aging. The retained austenite onase continually transformed during the first and second aging procedures. The decomposition of the retained austenite was accompanied by further precipitation of vanides [20], which reduced austenite stability, resulting in transformation from austenite to ferrite during the coling process. Fig. 7 also shows that, with increasing aging temperature, the fine carbides coalesced; as a result, the fine carbides tended to disappear when the samples were aged at 800°C. Because this process is controlled by diffusion, an increase in the aging temperature allowed sufficient time for the carbides to grow.

The dominant carbide in all of the double-aged samples was the  $M_6C$  carbide, followed by the  $M_7C_3$  carbide. The volume fraction of the  $M_{23}C_6$  carbides was smaller than that of the  $M_7C_3$  carbides. The different contrast of carbides in the SEM backscatter electron images was used to differentiate the type of carbide. In Fig. 8, the precipitates exhibited three different contrasts. The carbides with greater atomic number than that of the matrix exhibited white contrast

 $(M_6C)$ ; carbides that exhibited grey  $(M_{23}C_6)$  and lighter-grey  $(M_7C_3)$  contrast were also observed. The main metallic elements in the carbides were chromium, iron, vanadium, and tungsten. The SEM-EDS data for the dominant carbides  $M_6C$  and  $M_7C_3$  are given in Table 3. Notably, when the M<sub>7</sub>C<sub>3</sub> and M<sub>23</sub>C<sub>6</sub> carbides were analyzed at a spatial resolution of about 1 µm, interference occurred not only with the matrix but also with other carbide types. Because the volume fraction of the M23C6 carbides was small, only a few quantitative analyses were possible. The intensities of the chromium and iron peaks in the spectra of the M23C6 carbides were similar (the iron/chromium count ratio was about 1). The quantitative EDS data for the  $M_{23}C_6$  carbides showed that their average composition was 3.2%V-40.9%Cr-40.4%Fe-15.5%W (wt%). The EDS data showed that the M<sub>6</sub>C carbides were rich in tungsten, the M<sub>7</sub>C<sub>3</sub> carbides were rich in chromium and tungsten, and the M<sub>23</sub>C<sub>6</sub> and M<sub>7</sub>C<sub>3</sub> carbides were rich in chromium, with the chromium and tungsten contents in the M<sub>23</sub>C<sub>6</sub> carbides being respectively higher and lower than those in the M<sub>7</sub>C<sub>3</sub> carbide. Thus, the M<sub>7</sub>C<sub>3</sub> carbides exhibited a lighter grey contrast than the  $M_{23}C_6$  carbides. Notably, the  $M_6C$  and  $M_7C_3$  carbides were formed in the as-quenched microstructures; therefore, the  $M_6C$  and  $M_7C_3$  carbides in the double-aged sample were undissolved primary carbides and secondary carbides. The primary  $M_{23}C_6$  carbides dissolved during austenization and formed as secondary  $M_{23}C_6$  carbides during double-aging. The  $M_7C_3$  carbides were observed in two dif-

ferent sites; they coexisted with others carbide types, which indicated that they had formed via *in situ* transformation, and they were not joined with other carbide types, which indicated that they had formed individually in the matrix [21]. The  $M_6C$  carbides were present in all of the samples, and the  $M_{23}C_6$  carbides coexisted with the  $M_7C_3$  and  $M_6C$  carbides. These three carbide types had spherical and irregular morphologies.

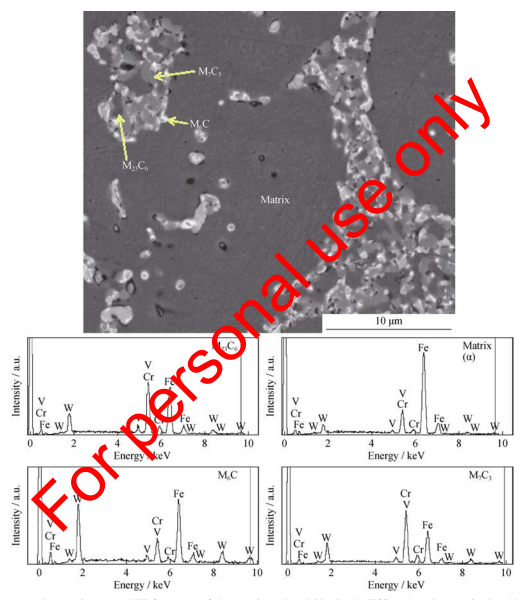


Fig. 8. Backscatter electron image and EDS spectra of the matrix and carbides in the H23 tool steel austenized at 1100°C and double-aged at 650°C.

Data in Table 3 would suggest that, at an austenizing temperature of 1100°C, the chromium content in the  $M_7C_3$  carbide increased at aging temperatures up to 750°C and then decreased significantly after aging at 800°C, whereas the tungsten content in the  $M_7C_3$  carbides increased with increasing aging temperature. In the sample aged at 650°C,

the  $M_7C_3$  carbides were very fine (Fig. 7), and as the aging temperature was increased to 750°C, these carbides grew and their composition moved closer to the equilibrium value. As the aging temperature was increased further to 800°C, the  $M_7C_3$  carbides began to lose their stability and their chromium contents decreased. Notably, the  $M_7C_3$  carbide is rich in chromium and the diffusivity of chromium is greater than those of tungsten and vanadium. Thus, after austenization at 1100°C, the M<sub>7</sub>C<sub>3</sub> carbides became unstable when aged at temperatures greater than 750°C. The chromium content in the M7C3 carbides was lower at an austenizing temperature of 1250°C than at an austenizing temperature of 1100°C when the same aging temperature was used, which indicated that the M<sub>7</sub>C<sub>3</sub> carbides in the sample austenized at the higher temperature were not stable at an aging temperature of 650°C. This finding is consistent with the results of a previous study [6] in which the authors reported that a higher austenizing temperature accelerated the carbides' approach to equilibrium. The evolution of the chemical composition of the M<sub>6</sub>C carbides was different than that of the  $M_7C_3$  carbides. The tungsten content of the  $M_6C$  carbides was higher at the austenizing temperature of 1250°C than at 1100°C when the aging temperature was the same. This higher tungsten content is attributed to the matrix becoming richer in tungsten at the higher austenizing temperature because more carbides had dissolved in it. The tungsten content in the M6C carbides in the double-tempered sample also increased with increasing aging temperature. When precipitated at 650°C, the secondary M<sub>6</sub>C carbides were fine and their composition was far from the equilibrium value. With

increasing aging temperature, the carbides grew by consuming more tungsten to approach the equilibrium composition. However, their tungsten content did not decrease after the steel was aged at 800°C, which indicated that these carbides still grew and did not reach equilibrium. Because the growth of carbides is controlled by diffusion (time and temperature dependence) and the diffusivity of tungsten is lower than that of the other alloying elements, more time and/or a higher temperature is required for the M6C carbides to become unstable. Fig. 9 shows a bright-field TEM image of MC, M<sub>6</sub>C, and M<sub>7</sub>C<sub>3</sub> carbides formed in the H23 tool steel double-tempered at 750°C; the TEM image does not show Fe<sub>3</sub>C or M<sub>2</sub>C carbides. These later carbides might have formed in the early stages of transformation and dissolved quickly with increasing aging temperature. Baker and Nutting [22] also studied the propitation sequence of carbides in ferritic microstructures in a 2.25%Cr-1%Mo (wt%) steel. The ferrite contract  $M_2C$  carbides. Subsequent aging led to the dissolution on the M<sub>2</sub>C and its replacement by the more stable Mc earb les. In the present work, the M7C3 carbides nucleated by tween pre-existing M6C carbides or the M6C orbides acted as nucleation sites for M<sub>7</sub>C<sub>3</sub> secondary caride. Fine MC carbides were observed in the matrix; howev r, no evidence of the  $M_{23}C_6$  carbides was observed.

Table 3.         SEM-EDS results or an ides in the double-aged H23 tool store						wt%
$T_{\gamma} / ^{\circ}\mathrm{C}$	$T_{\mathrm{DA}} / ^{\circ}\mathrm{C}$		Comment*			
$I_{\gamma} / C$		Fe	Cr Cr	V	W	Comment
	650	54.6 3	$11.8\pm0.3$	$0.9\pm0.1$	$32.7 \pm 3.2$	
	750	10.1+ J.J	$11.9\pm2.6$	$2.6\pm0.1$	$36.5\pm2.0$	M <sub>6</sub> C
1100 -	800	3, 5 = 5.2	$11.8 \pm 1.6$	$1.9\pm0.1$	$46.9\pm2.6$	
1100	650	44.8±3.6	$35.2 \pm 4.0$	$3.1\pm0.1$	$16.9\pm0.4$	
	750	$34.5 \pm 0.1$	$38.8\pm0.1$	$3.7\pm0.1$	23.0±0.2	$M_7C_3$
	<b>97</b> 0	$60.7\pm6.4$	$13.2 \pm 1.1$	$1.3\pm0.6$	$24.8\pm4.7$	
	650	37.1 + 5.3	$11.8 \pm 1.2$	$2.4\pm0.2$	$48.7\pm5.9$	
	750	$28.1\pm2.2$	$8.3\pm0.1$	$2.6\pm0.2$	$61.0 \pm 2.1$	M <sub>6</sub> C
1250 -	800	$25.6\pm2.4$	$8.7\pm0.4$	$2.5\pm0.2$	$63.1 \pm 2.8$	
1230	650	$46.4\pm1.8$	$19.6\pm0.8$	$2.2\pm0.1$	$31.7\pm1.0$	
	750	$70.5\pm2.5$	$16.3\pm2.6$	$1.4\pm0.2$	$11.9\pm0.5$	$M_7C_3$
	800	$77.2 \pm 2.1$	$12.3 \pm 1.3$	$1.0 \pm 0.1$	$9.5\pm0.3$	

Note: <sup>\*</sup> The quantitative data should be considered with caution given the size of carbides;  $T_{DA}$  is the double-aging temperature.

#### 3.3.2. Carbide evolution

Four types of carbides were observed in the double-aged H23 tool steel: MC,  $M_6C$ ,  $M_7C_3$ , and  $M_{23}C_6$  carbides. The dominant carbides formed under the double-aging condi-

tions were the  $M_6C$  and  $M_7C_3$  carbides. The formation of the  $M_6C$  carbide at the expense of the  $M_{23}C_6$  carbide was not observed. Secondary  $M_6C$  carbides formed within the tung-sten-rich matrix via dissolution of the  $M_2C$  carbides. Al-

though the Fe<sub>3</sub>C carbides were not observed in all of the double-aged microstructures, we hypothesized that this type of carbide precipitated below the aging temperature of 650°C and transformed quickly into the chromium-rich carbides  $M_7C_3$  and  $M_{23}C_6$ . The  $M_{23}C_6$  carbides grew with increasing aging temperature and gradually disappeared from the microstructure after the sample was aged at 800°C. The  $M_7C_3$  carbide either formed via the reaction  $M_3C \rightarrow M_7C_3 + M_{23}C_6$  or nucleated directly from ferrite (separate or individual nucleation). In the former case, the reaction started by the increase of the chromium content in the  $M_3C$  carbide and the  $M_7C_3$  carbide particles grew. The  $M_7C_3$  carbide was more dominant than the  $M_{23}C_6$  carbide in the double-aged

H23 tool steel because of the presence of vanadium in the steel, which tends to stabilize the  $M_7C_3$  carbide phase [23]. Bjärbo and Hättestrand [24] have reported that a competition occurs between the nucleation and growth processes of the  $M_7C_3$  and  $M_{23}C_6$  carbides, and in high-chromium-content tool steel, the  $M_{23}C_6$  carbide would nucleate at the same time as the  $M_7C_3$  carbide. We concluded that the Fe<sub>3</sub>C transformed into  $M_7C_3$  carbide, and a small volume fraction of  $M_3C$  likely transformed into  $M_{23}C_6$  carbide. Therefore, the carbide evolution in the H23 tool steel occurred as follows. *in situ* transformation: Matrix  $\rightarrow M_3C \rightarrow M_7C_3 + M_{23}C_6$ ; separate transformation: Matrix  $\rightarrow M_3C \rightarrow M_2C \rightarrow M_6C$  and Matrix  $\rightarrow M_7C_3$ .

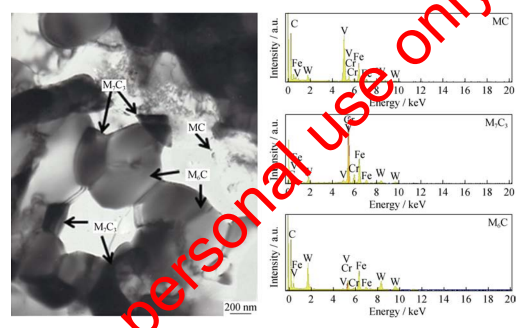


Fig. 9. Bright-field TEM image of the h.3 tool steel water quenched from 1100°C and double-aged at 750°C and EDS spectra of carbides.

The XRD diffratograms in Fig. 10 confirmed the presence of solid-solution ferrite ( $\alpha$ -Fe–Cr), the M<sub>6</sub>C, M<sub>23</sub>C<sub>6</sub>, and M<sub>7</sub>C<sub>3</sub> carbides. The MC carbide was not identified by XRD; however, its presence was confirmed by TEM [10]. The low volume fraction of the MC carbide and the overlap of its peaks with those of the M<sub>7</sub>C<sub>3</sub> and M<sub>23</sub>C<sub>6</sub> carbides [25] precluded its detection by XRD. Fig. 10 also shows that the XRD peaks of the M<sub>6</sub>C carbide were dominant and the intensity of the XRD peaks of the M<sub>7</sub>C<sub>3</sub> carbide was higher than that of the M<sub>23</sub>C<sub>6</sub> carbide under all double-aging conditions. The XRD results were in good agreement with the other microstructure studies regarding the high volume fraction of the M<sub>6</sub>C carbides and the dominance of the M<sub>7</sub>C<sub>3</sub> over the M<sub>23</sub>C<sub>6</sub> carbides.

3.3.3. Effect of double-aging treatment on hardness

The precipitation of carbides during aging significantly contributed to the final hardness of the tool steel. The effects of the austenizing and double-aging temperatures on the hardness of the H23 tool steel are shown in Fig. 11. Fig. 11 shows that the hardness of the double aged H23 tool steel increased with increasing austenizing temperature and decreased with increasing double-aging temperature. The highest hardness (448 HV) was achieved in the case of the sample austenized at 1250°C and double aged at 650°C. This result is attributable to the carbides dissolving more extensively into the matrix with increasing austenizing temperature, thereby providing a high nucleation rate of carbide precipitation during the double-aging process [6]. Double-

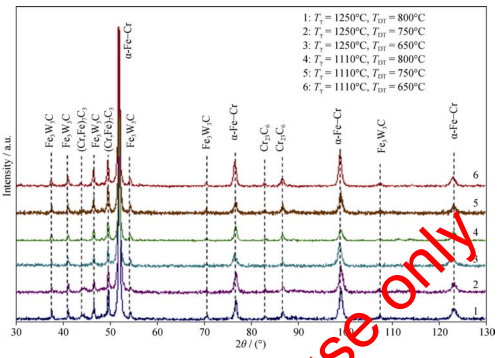


Fig. 10. XRD diffraction pattern of the H23 tool steel with different austenization and double-aging temperatures. The ICDD cards used were α-Fe-Cr (34–396), austenite (52–512), Fe<sub>3</sub>W<sub>3</sub>C (41–1351), Cr<sub>23</sub>C<sub>6</sub> (35–7)<sup>3</sup>), and (Cr,Fe)<sub>7</sub>C<sub>3</sub> (5–720).

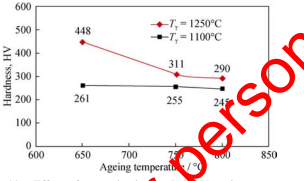


Fig. 11. Effect of austenization and double-aging temperatures on the hardness of the H22 to 1 steel.

aging at temperatures of  $6.9^{\circ}$ C and  $800^{\circ}$ C did not reveal a secondary hardening teak, which has been correlated to fine M<sub>2</sub>C carbides [1]. In the microstructure of the double-aged H23 tool steel, unstable M<sub>2</sub>C or M<sub>3</sub>C carbides were not observed. The higher chromium content in the H23 tool steel caused these unstable carbides to dissolve at temperatures below 650°C. Quantitative data of carbide size and volume fraction for the three double-aging temperatures are given in rate 4.

Table 4 shows that the volume fraction of carbides slightly increased with increasing aging temperature, which was attributed to the precipitation of secondary carbides; more of these carbides precipitated at the higher aging temperature because of the greater diffusivity of the alloying elements. The secondary carbides formed during the double-aging process of the H23 tool steel were M6C and chromium-rich carbides M7C3 and M23C6. The latter carbides reached their equilibrium composition and dissolved at lower temperature compared to the M<sub>6</sub>C carbide; in addition, increasing austenizing temperature accelerated the approach of the chromium-rich carbides to equilibrium during aging at 650°C. Hence, no significant difference in the volume fraction of carbides was observed between steels aged at 750°C and 800°C. In addition, no correlation was observed between the average size of carbides and aging temperature, likely because coalesced carbides formed networks that

Table 4. Volume fraction and average size of carbides in the H23 tool steel austenized and double-aged at various temperatures

<u> </u>	Double ageing temperature / °C						
$T_{\gamma} / ^{\circ}\mathrm{C}$	650		750		800		
	V <sub>c</sub> / %	Mean size / $\mu m$	V <sub>c</sub> / %	Mean size / $\mu m$	V <sub>c</sub> / %	Mean size / $\mu m$	
1100	$16.0\pm0.2$	$0.3\pm0.1$	$17.7\pm0.5$	$0.2\pm0.2$	$18.2\pm0.3$	$0.3\pm0.2$	
1250	$16.9\pm0.3$	$0.3 \pm 0.1$	$18.3\pm0.6$	$0.3 \pm 0.1$	$18.5\pm0.4$	$0.3\pm0.2$	

made the measurements of carbide size less accurate. The average carbide size in the double-aged H23 tool steel increased with increasing aging temperature when the steel was austenized at 1250°C because of a reduction in the total area of the ferrite/carbide interphase boundary. The overall impression from the microstructure studies was that carbides were slightly coarsened with increasing aging temperature. We concluded that the decrease of hardness with increasing aging temperature was due to the carbides coarsening and ferrite grain growth. This conclusion suggests that the peak of secondary hardening of the H23 tool steel occurred at temperatures below 650°C.

#### 3.4. Equilibrium microstructure at 600°C

The calculated isopleth phase diagram is shown in Fig. 12, where the vertical dotted line indicates the composition of the steel used in this study. According to the phase diagram, the equilibrium microstructure of this steel at 600°C should consist of ferrite,  $M_{23}C_6$  carbide, and a Fe<sub>2</sub>W Laves phase. To verify the thermodynamic calculations, the H23 tool steel was austenized at 1250°C for 1 h, water quenched, and then heat treated at 600°C for 400 h. According to the XRD data shown in Fig. 13, the phases present in the microstructure were ferrite and the  $M_6C$  and  $M_{23}C_6$  carbides. No peaks corresponding to the Fe<sub>2</sub>W Laves phase were observed, which we attributed either to the absence of this phase in the microstructure of the steel or its presence at a volume fraction below the detection limit of the X-ray diffractometer. Given the XRD results, the experimental microstructure and the results of the thermodynamic calculation did not agree with each other.

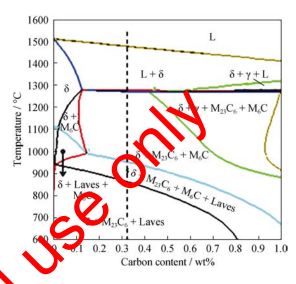


Fig. 12. Calculated isopleth of the H23 tool steel phase diarr n [10].

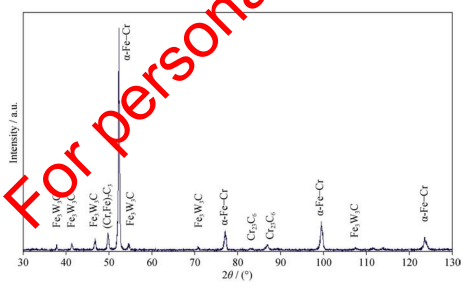


Fig. 13. XRD diffraction pattern of the H23 tool steel water quenched from  $T_{\gamma} = 1250^{\circ}$ C and single aging at 600°C for 400 h. The ICDD cards used were those of  $\alpha$ -Fe-Cr (34–396), austenite (52–512), Fe<sub>3</sub>W<sub>3</sub>C (41–1351), Cr<sub>23</sub>C<sub>6</sub> (35–783), and (Cr,Fe)<sub>7</sub>C<sub>3</sub> (5–720).

# 4. Conclusions

In this paper, the effect of austenization and double-aging treatments on the microstructure and hardness of H23 tool steel were studied. The following conclusions were drawn:

(1) The  $M_6C$  (Fe<sub>3</sub>W<sub>3</sub>C),  $M_7C_3$  ((Cr,Fe)<sub>7</sub>C<sub>3</sub>), and MC carbides were formed in the as-quenched steels austenized at

#### 1100°C and 1250°C.

(2) The chromium content of the  $M_7C_3$  carbides tended to decrease at higher austenizing and aging temperatures, and the tungsten content of the  $M_6C$  carbides increased with increasing austenizing and aging temperatures.

(3) The MC,  $M_6C$ ,  $M_7C_3$ , and  $M_{23}C_6$  carbides were observed in the microstructure after the double-aging treatment,

#### Int. J. Miner. Metall. Mater., Vol. 22, No. 3, Mar. 2015

and M<sub>6</sub>C and M<sub>7</sub>C<sub>3</sub> were the dominant carbides.

(4) The secondary  $M_6C$  carbides formed during double-aging because of the dissolution of  $M_2C$  carbides.

(5) The approach of carbides to their equilibrium composition during double-aging was strongly affected by the austenizing temperature.

(6) The maximum hardness of the double-aged H23 tool steel (448 HV) was achieved via austenization at 1250°C followed by double-aging at 650°C.

# Acknowledgements

The author (MN) would like to express great appreciation to the Directorate of Higher Education, Indonesian Government and to the Institut Teknologi Nasional (Itenas), Bandung, Indonesia for their financial support.

### References

- G.A. Roberts, G. Kraus, and R.L. Kennedy, *Tool Steels*, Materials Park, ASM International, Ohio, 1998.
- [2] M.L. Fares, M. Athmani, Y. Khelfaoui, and A. Khettache, An investigation into the effects of conventional heat treatments on mechanical characteristics of new hot working tool steel, *IOP Conf. Ser. Mater. Sci. Eng.*, 28(2012), No. 1, art. N. 012024.
- P. Bała and J. Krawczyk, Transformation during quinching and ageing of hot-work tool steel, *Hradec Moravic*, 190209 p. 1.
- [4] Z. Zhang, D. Delagnes, and G. Bernhart, Micro tructure evolution of hot-work tool steels during tem ering and definition of a kinetic law based on hardness measurements, *Mater. Sci. Eng. A*, 380(2004), p. 222.
- [5] A. Medvedeva, J. Bergström S. Gunnarsson, and J. Andersson, High temperature preparties and microstructural stability of hot work tool needs, *Mate. Sci. Eng. A*, 523(2009), No. 1-2, p. 39.
- [6] M. Nurbanasari, P. Tsakiropoulos, and E.J. Palmiere, A study of carbide precipitation in a H21 tool steel, *ISLJ Int.*, 54(2014), No. 7, p. 1667.
- [7] R.C. Thomson and H.K.D.H. Bhadeshia, Carbide precipitation in 12Cr1MoV power plant steel, *Metall. Trans. A*, 23(1992), p. 1171.
- [8] N. Dudova and R. Kaibyshev, On the precipitation sequence in a 10% Cr steel under tempering, *ISLJ Int.*, 51(2011), No. 5, p. 826.
- [9] M.A. Asadabad, S. Kheirandish, and A.J. Novinrooz, Tem-

pering behavior of 4.5Cr-2W-0.25V steel, J. Iron Steel Res. Int., 17(2010), No. 10, p. 57.

- [10] M. Nurbanasari, P. Tsakiropoulos, and E.J. Palmiere, On the solidification of a H23 tool steel, *Trans. Indian Inst. Met.*, 67(2014), No. 6, p. 935.
- [11] N. Mebarki, D. Delagnes, P. Lamesle, F. Delmas, and C. Levaillant, Relationship between microstructure and mechanical properties of a 5% Cr tempered martensitic tool steel, *Mater. Sci. Eng. A*, 387-389(2004), p. 171.
- [12] A.M. Elrakayby and B. Mills, Identification of carbides in high-speed steels, J. Mater. Sci. Lett., 5(1986), p. 332.
- [13] F. Abe, H. Araki, and T. Noda, The effect of tungsten on dislocation recovery and precipitation behavior of low-activation martensitic 9Cr steels, *Metall Trans.* 4, 22(1991), p. 2225.
  [14] F.B. Pickering, The properties of toor teels for mould and die
- [14] F.B. Pickering, The properties of toor teels for mould and die applications [in] G. Krauss and H. Nordberg, eds., *Tool Materials for Molds and Dats: Application and Performance*, CSM Press Golden, Color do, 1987, p. 3.
- [15] D.J. Dyson and K.W. Andrews, Carbide M<sub>7</sub>C<sub>3</sub> and its formation in alloyste is *J. Iron Steel Inst.*, 207(1969), No.2, p.208.
- [16] K. Kuo care to in chromium, molybdenum, and tungsten steels *5. nrop* Steel Inst., 173(1953), p. 363.
- [17] P. Wilson, Metallurgy and Heat Treatment of Tool Steel, Mc Grave Jill, London, 1975.
- 18, G.E. Totten, L. Xie, and K. Funatani, *Handbook of Mechanical Alloy Design*, Marcel Dekker Inc, New York, 2004.
- S. Karagoz and H. Fischmeister, Microstructure and toughness in high speed tool steels: the influence of hot reduction and austenitization temperature, *Steel Res.*, 58(1987), No. 8, p. 353.
- [20] A. Saha Podder and H.K.D.H. Bhadeshia, Thermal stability of austenite retained in bainitic steels, *Mater. Sci. Eng. A*, 527(2010), No. 7-8, p. 2121.
- [21] G.J. Cai, H.O. Andren, and L.E. Svensson, Microstructural change of a 5% Cr steel weld metal during tempering, *Mater. Sci. Eng. A*, 242(1998), No. 1-2, p. 202.
- [22] R.G. Baker and J. Nutting, The tempering of 2.25Cr%–1%Mo steel after quenching and normalizing, *J. Iron Steel Inst.*, 202 (1959), p. 257.
- [23] H.K.D.H. Bhadeshia and R.W.K. Honeycombe, *Steels: Mi-crostructure and Properties*, Elsevier Ltd., Oxford, 2006.
- [24] A. Bjärbo and M. Hättestrand, Complex carbide growth, dissolution and coarsening in a modified 12 pct chromium steel-an experimental and theoretical study, *Metall. Mater. Trans. A*, 32 (2001), No. 1, p. 19.
- [25] K. Asakura, A. Kohyama, and T. Yamada, Mechanical properties and microstructure changes of low-activation 3Cr-2W-V-Ti ferritic steels developed for nuclear applications, *ISIJ Int.*, 30(1990), No. 11, p. 947.

#### 284