

## Article

# Influence of Vacuum Heat Treatments on Microstructure and Mechanical Properties of M35 High Speed Steel

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**Abstract:** Towards the end of the last century, vacuum heat treatment of high speed steels was increasingly used in the fabrication of precision cutting tools. This study investigates the influence of vacuum heat treatments at different pressures of quenching gas on the microstructure and mechanical properties of taps made of M35 high speed steel. Taps were characterized by optical microscopy, scanning electron microscopy with energy dispersive spectroscopy, X-ray diffraction, apparent grain size and Vickers hardness measurements, and scratch tests. Failure analysis after tapping tests was also performed to determine the main fracture mechanisms. For all taps, the results showed that microstructures and the values of characteristics of secondary carbides, retained austenite, apparent grain size and Vickers hardness were comparable to previously reported ones for vacuum heat treated high speed steels. For taps vacuum heat treated at six bar, the highest plane strain fracture toughness was due to a higher content of finer small secondary carbides. In contrast, the lowest plane strain fracture toughness of taps vacuum heat treated at eight bar may be due to an excessive amount of finer small secondary carbides, which may provide a preferential path for crack propagation. Finally, the predominant fracture mechanism of taps was quasi-cleavage.

Keywords: high speed steel; vacuum heat treatment; microstructure; plane strain fracture toughness

## 1. Introduction

High speed steels (HSS) are a complex class of tool steels continuously improved for applications in the global cutting tool market as wear-resistant materials for drills, taps, milling cutters, broaches, slotting tool and hobs. These steels must withstand abrasive or adhesive wear and have sufficient hardness, toughness and ductility to prevent chipping, galling, cracking, etc. [1].

In the fabrication of precision cutting tools, heat treatment of HSS in a salt bath has been replaced by vacuum heat treatment with uniform high-pressure gas quenching, since this procedure avoids waste and the need for washing the heat treated tools. A typical vacuum heat treatment includes several pre-heat, high heat, quenching and tempering stages. The final microstructure consists of large primary carbides in a tempered martensite matrix, hardened by the precipitation of uniformly distributed secondary carbides [2]. Nevertheless, the vacuum heat treatment has its limitations, especially concerning the number of pre-heats and holding time at high heat. Moreover, steel suppliers recommend only general guidelines about the quench rate of many grades of HSS in vacuum heat treatment [3].



The role of microstructural features is well-known in enhancing the mechanical properties of steels [2,4,5]. Over the past few decades, many studies have been carried out on wear resistance and performance of high speed steels after vacuum heat treatment. The effects of different austenitizing and tempering temperatures were evaluated, and the microstructure was optimized for increasing fracture toughness, hardness, fatigue and wear resistance of the tool [6–11]. With respect to fracture toughness, several authors used circumferentially notched and fatigue pre-cracked tensile test specimens, according to previously published methods [12,13]. The advantage of such specimens is their radial symmetry, which makes them specifically suitable for studying the influence of the microstructure on fracture toughness of metals. Due to the radial symmetry of heat transfer during heat treatment, the microstructure forming along a circumferential area is completely uniform. Moreover, the fatigue pre-crack can be created before the heat treatment without detrimental effects on the crack tip blunting. However, this method is not only difficult to apply, but also time consuming and expensive [14]. The most common alternative method is the Vickers indentation fracture test, where the fracture toughness is determined through a Vickers probe and by expressions accounting for indentation load, Young's modulus, hardness, residual stresses, plastic dissipation inside the material and the nature of cracks produced during indentation [15–17]. Recently, Sola et al. proposed a novel technique to measure the fracture toughness of tool steels by scratch test [18]. Generally, the scratch test consists of pulling a probe across the surface of the material under a controllable applied load. According to the authors, the failure mode (fracture or plastic yielding) depends on material properties and geometry of the scratching tool. It is therefore possible to link the forces acting on the scratch tip and on the tool geometry to the plane strain fracture toughness ( $K_C$ ) by an equation accounting for the tangential force necessary to move the indenter, the indenter tip width and the measured penetration depth [16].

The present study investigates microstructure and mechanical properties of taps made of M35 high speed steel and vacuum heat treated at increasing pressures of quenching gas for improving their fracture toughness and durability. For this purpose, the microstructure and plane strain fracture toughness of taps undergoing conventional vacuum heat treatment were characterized by optical microscopy (OM), scanning electron microscopy with energy dispersive spectroscopy (SEM/EDS), X-ray diffraction (XRD), apparent grain size and Vickers hardness measurements, and scratch tests. Tapping tests and failure analyses were also performed on the same taps to determine the main fracture mechanisms involved during normal operating conditions. Finally, a new set of taps was vacuum heat treated at increasing pressures of quenching gas, and the resulting microstructure and plane strain fracture toughness were evaluated by the above mentioned techniques.

#### 2. Materials and Methods

#### 2.1. Chemical Composition and Conventional Heat Treatment of Taps

The material used in this investigation was commercially available M35 high speed steel with the following chemical composition (wt.%): 0.935% C, 0.362% Si, 0.233% Mn, 0.012% P, 0.001% S, 5.018% Co, 3.803% Cr, 4.643% Mo, 1.830% V, 6.613% W, balance Fe. The material was shaped in the form of cylindrical taps 10 mm in diameter and 100 mm in length.

The conventional heat treatment (indicated as CHT\_5) of taps was carried out in a horizontal vacuum tube furnace with uniform high-pressure gas quenching, using nitrogen gas at a pressure of 5 bar and quenching speed of 10 °C/s. Quenching gas flew through nozzles located circumferentially on the walls of the furnace. After three preheat stages at 550, 850 and 1000 °C, the taps were heated at 2 °C/min to the austenitizing temperature of 1195 °C and maintained at this temperature for 10 min. Taps were then gas quenched from the austenitizing temperature to a temperature of 30 °C. Finally, triple tempering at 560 °C for 2 h were performed in the same furnace. All heat treatment parameters were selected on the basis of the recommendations of the steel supplier. The scheme in Figure 1 shows the CHT\_5 treatment applied to the taps.



Figure 1. Conventional heat treatment (CHT\_5) applied to the taps.

#### 2.2. Material Characterization

Material characterization was performed on taps pulled out of the horizontal vacuum furnace after quenching and tempering. To determine the microstructure of alloy, longitudinal sections (parallel to the metal surface) and cross-sections (perpendicular to the metal surface) of the taps were prepared, mounted in resin, polished and examined after chemical etching by Picral (4 g picric acid in 100 mL ethanol) through a Leica MEF4M optical microscope (Leica, Wetzlar, Germany) and a Zeiss EVO MA 15 (Zeiss, Oberkochen, Germany) scanning electron microscope (SEM), equipped with an Oxford X-Max 50 (Oxford Instruments, Abingdon-on-Thames, UK) energy dispersive microprobe for semi-quantitative analyses (EDS). The SEM micrographs were recorded in secondary electron imaging (SEI-SEM) and back-scattered electron (BSE-SEM) mode. The carbide particles visible in the matrix were classified as primary carbides (PCs: size > 5.0  $\mu$ m) and secondary carbides (SCs: size  $\leq$  5.0  $\mu$ m). The secondary carbides were further sub-classified as large secondary carbides (LSCs:  $1.0 \ \mu m < size \le 5.0 \ \mu m$ ) and small secondary carbides (SSCs:  $0.1 \,\mu\text{m} \le \text{size} \le 1.0 \,\mu\text{m}$ ) [19]. The micrographs of the cross-sections after tempering were processed by Leica Application Suite (LAS V4.12.0, Leica, Wetzlar, Germany) image analysis software to evaluate some characteristics of large and small secondary carbides, such as area fraction, population density and size, according to a previously described procedure [20]. A mean of 1000 secondary carbides was considered for each tap to achieve appropriate statistical reliability associated with these measurements.

On the cross-sections after tempering, the apparent grain size was determined in agreement with the EN ISO 643:2020 standard [21], whereas the type of carbide particles and retained austenite content (RA) were evaluated by X-ray diffraction (XRD) analyses of the bulk taps with a Philips X'PERT PW3050 diffractometer (Philips, Amsterdam, The Netherlands), using Cu-K $\alpha$  radiations and diffracted beam monochromator, with an intensity scanner versus diffraction angle between 38° and 105° (0.01° step size, 2 s/step scanner velocity and 1.5 grid), a 40 kV voltage and a 40 mA filament current. The calculation of RA was performed according to the ASTM E975-13 standard [22].

The Vickers hardness measurements under 1000  $g_f$  load and 15 s loading time (HV1) were carried out in triplicate on the cross-sections by a Future-Tech FM-110 Vickers microindenter (Future-Tech Corp., Kawasaki, Japan). The Vickers hardness values were then converted in Rockwell C hardness values (HRC) in agreement with the ASTM E140-12B standard [23].

Finally, the fracture toughness was determined by scratch tests on the cross-sections after tempering and in agreement with a previously published method [18,24,25]. Prior to the tests, the surfaces of the samples were prepared according to a previously published procedure [26]. The scratch test produced a scratch in the material using a probe drawn across the sample under a constant vertical load and with constant speed. Compared to the standard fracture toughness tests, this technique provided the following advantages: (i) It was non-destructive; (ii) it did not require specific dimensions of the

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samples, nor an initial notch, pre-crack or fatigue crack; (iii) it allowed the sampling of the fracture properties at different locations in the material; (iv) it had a minimal cost and (v) could be performed on a limited supply of the material. The scratch tests were carried out by a Revetest Scratch Tester (CSM Instrument, Needham, MA, USA), using a 200  $\mu$ m Rockwell C diamond probe, a scratching speed of 6 mm/min, an applied load of 100 N and a scratch length of 6 mm. In agreement with a previously published method [25], the plane strain fracture toughness K<sub>C</sub> was determined by the following equation,

$$K_C = \frac{F_t}{\sqrt{2wd(w+2d)}} \tag{1}$$

where  $F_t$  is the tangential force necessary to move the indenter (measured by the equipment), w is the indenter tip width and d is the penetration depth. The penetration depth was directly calculated by the scratch tester. The equation was derived from previous studies on fracture mechanics framework [25,26].

#### 2.3. Tapping Tests

Tapping tests were performed by a TH6563x63 (Shenyang Machine Tool Co., Ltd., Liaoning, China) horizontal tester machine. A grounded workpiece in 30CrMnTi steel,  $700 \times 500 \times 48 \text{ mm}^3$  in volume and with hardness equal to 210 HBW 2.5/187.5/30 was inserted and fixed inside the machine. Holes arranged in rows were made inside the workpiece. Taps were then rotated and moved into holes with a cutting speed of 7.5 m/min. A cutting fluid based on an emulsion of mineral oil at 5% was applied at a rate of 5 L/min. The criterion for end of tap life was its catastrophic failure. A scheme of the tapping test is shown in Figure 2.



Figure 2. Scheme of the tapping test.

#### 2.4. Failure Analysis

After tapping tests, fracture surfaces of taps were first observed by a Leica MZ6 (Leica) stereomicroscope. To identify the main fracture mechanisms, morphological observations were performed by the Zeiss EVO MA 15 (Zeiss) scanning electron microscope with the Oxford X-Max 50 (Oxford Instruments) energy dispersive microprobe. Metallographic analyses of properly polished cross-sections of the fracture surfaces were conducted after chemical etching by Picral under the Leica MEF4M optical microscope (Leica).

#### 2.5. Vacuum Heat Treatments

A new set of taps with the same chemical composition of those in Section 2.1 was heat treated in the horizontal vacuum furnace according to the treatment schematized in Figure 1, but using nitrogen gas at pressures of 6 and 8 bar (indicated as CHT\_6 and CHT\_8, respectively). During each heat treatment, the material was characterized following the procedure described in Section 2.2.

#### 3. Results

#### 3.1. Microstructural Characterization and Hardness Variation

Optical and BSE-SEM micrographs of the microstructure observed on the cross-section of taps, after CHT\_5 heat treatment, are shown in Figure 3. The microstructure exhibited a non-uniform distribution of large and dendritic-type primary carbides (marked by PCs), and a fairly uniform distribution of secondary carbides in a tempered martensite matrix, with retained austenite and resolved prior austenite grain boundaries. The two types of secondary carbides, classified according to their size, appeared either as well-defined white regions (marked by LSCs) or tiny white patches (marked by SSCs) (Figure 3a). In Figure 3b, a semi-continuous distribution of proeutectoid carbides was also visible along prior austenitic grain boundaries. The mean value of area fraction of large secondary carbides measured on taps cross-sections after CHT\_5 heat treatment was  $2.76 \pm 0.23\%$ , whereas those of population density and size were  $4.0 \pm 0.5 \times 10^3/\text{mm}^2$  and  $2.78 \pm 0.12 \,\mu\text{m}$ , respectively. The mean value of area fraction of small secondary carbides measured on the same regions was  $2.53 \pm 0.15\%$ , while that of population density was  $134 \pm 4 \times 10^3/\text{mm}^2$ . The mean value of size of small secondary carbides measured on the same regions was  $0.74 \pm 0.08 \,\mu\text{m}$ .



**Figure 3.** (**a**) Optical and (**b**) BSE-SEM micrographs of the microstructure observed on the cross-section of taps after CHT\_5. Yellow circles in (**a**) indicate primary carbides (PCs), large secondary carbides (LSCs) and small secondary carbides (SSCs). Yellow arrows in (**b**) show the semi-continuous distribution of proeutectoid carbides visible along prior austenitic grain boundaries.

Comparing the EDS semi-quantitative analyses of carbide particles (Figure 4), after quenching the primary carbides were rich in V or in W and Mo (spectra 1 and 2 in Figure 4), and changed only marginally their chemical composition after triple tempering at 560 °C. After quenching, secondary carbides and proeutectoid carbides were rich in Fe and poor in Cr, and maintained their chemical composition until the end of the heat treatment (spectrum 3 in Figure 4). Nevertheless, the EDS results proved that during tempering there was a precipitation of new Cr-rich secondary carbides. The XRD diffractograms recorded on the bulk taps after CHT\_5 heat treatment revealed the presence of martensite (indexed as ferrite, since the resolution of the diffractometer was not sufficient to resolve the

tetragonality of martensite), alloyed cementite (indexed as Fe<sub>3</sub>C), MC, M<sub>6</sub>C, M<sub>7</sub>C<sub>3</sub> and RA (Figure 5). The mean value of RA in the matrix measured by XRD was  $12.4 \pm 0.5\%$ .



**Figure 4.** (a) BSE-SEM micrograph of the microstructure observed on the cross-section of taps after quenching and (b) semi-quantitative EDS analyses (wt.%) of different carbide particles in the same microstructure.



Figure 5. XRD diffractogram recorded on the bulk taps after CHT\_5.

The mean value of apparent grain size measured on tap cross-sections after CHT\_5 heat treatment was 2.6  $\pm$  0.1  $\mu$ m (G14, according to the EN ISO 643:2020 standard), whereas that of plane strain fracture toughness, calculated by Equation (1), was 31  $\pm$  5 MPa·m<sup>1/2</sup>. After quenching, the mean value

of Vickers hardness measured on the same regions was 914 HV1 (corresponding to 67 HRC). Triple tempering at 560 °C resulted in Vickers hardness of 892 HV1, corresponding to 66 HRC.

#### 3.2. Failure Analysis after Tapping Tests

After tapping tests, the stereomicroscopy images of the fracture surfaces of taps had a smooth and flat appearance, typical of brittle materials. At higher magnification, the SEI-SEM analyses revealed that the morphology of the fracture surfaces (Figure 6a) was characterized by two features: classical cleavage fracture of primary carbides (marked as C in Figure 6b) and quasi-cleavage fracture of the matrix (marked as the yellow rectangle QC in Figure 6b). In the first case, cracking of primary carbides was visible, and microcracks occurred at the interface between the primary carbide and the matrix. The EDS semi-quantitative analyses on the cleavage regions of the fracture surfaces confirmed that these regions corresponded to primary carbides. In the second case, a large number of microvoids were formed by decohesion of secondary carbides (yellow arrows in Figure 6c). No cracking of secondary carbides was detected. The optical micrograph of the fracture surfaces observed on the cross-sections of taps is shown in Figure 6d. These surfaces were mostly irregular. Fractures were deflected and partially surrounded the numerous secondary carbides detected close to the fracture surfaces, also suggesting that these carbide particles provided a preferential path for crack propagation.





**Figure 6.** (**a**–**c**) SEI-SEM micrographs of the fracture surfaces of taps after tapping test showing, in the order, an overview of fracture surface, details of cleavage (C) and a quasi-cleavage (QC, yellow rectangle) region, and microvoids formed by decohesion of secondary carbides (yellow arrows); (**d**) optical micrograph of the fracture surfaces observed on the cross-sections of taps.

#### 3.3. Influence of Vacuum Heat Treatments on Microstructure and Fracture Toughness

Optical and scanning electron micrographs of the microstructure observed on the cross-sections of taps after CHT\_6 and CHT\_8 heat treatments (Figure 7a,c, respectively) were similar to those after CHT\_5 (Figure 3a), but the extent of the semi-continuous distribution of proeutectoid carbides along prior austenitic grain boundaries decreased with increasing pressure of quenching gas (Figure 7b,d). During CHT\_6 and CHT\_8 heat treatments, primary and secondary carbides, detected by EDS and XRD, were unchanged with respect to those observed after CHT\_5.



**Figure 7.** (**a**,**c**) Optical and (**b**,**d**) BSE-SEM micrographs of the microstructure observed on the cross-section of taps after CHT\_6 and CHT\_8 heat treatments. Yellow arrows in (**b**,**d**) show the semi-continuous distribution of proeutectoid carbides visible along prior austenitic grain boundaries.

The characteristics of large and small secondary carbides at increasing pressure of quenching gas are shown in Figure 8a–c. The mean values of area fraction (Figure 8a) and population density (Figure 8b) of large and small secondary carbides progressively increased with increasing pressure of quenching gas. The mean value of area fraction of large secondary carbides increased up to the final value of  $3.21 \pm 0.19\%$  (corresponding to an overall increase of about 16%), while that of small secondary carbides reached the final value of  $3.49 \pm 0.17\%$  (corresponding to an overall increase of about 38%). The mean value of population density of large secondary carbides raised up to the final value of  $9.0 \pm 0.3 \times 10^3$ /mm<sup>2</sup> (corresponding to an overall increase of about 125%), whereas that of small secondary carbides reached the final value of  $167 \pm 3 \times 10^3$ /mm<sup>2</sup> (corresponding to an overall increase of about 25%). Conversely, the mean value of size of the same carbide particles was reduced with an increase in pressure of quenching gas (Figure 8c). The mean value of size of large secondary carbides decreased up to the final value of  $2.17 \pm 0.18 \,\mu$ m (corresponding to an overall decrease of about 22%), while that of small secondary carbides reached the final value of  $2.17 \pm 0.18 \,\mu$ m (corresponding to an overall decrease of about 22%).



**Figure 8.** Characteristics of large and small secondary carbides at increasing pressure of quenching gas: mean values of (**a**) area fraction, (**b**) population density and (**c**) size.

The mean values of retained austenite, apparent grain size, plane strain fracture toughness and Vickers hardness of taps after the different heat treatments are reported in Table 1. The mean value of RA decreased with increasing pressure of quenching gas, reaching the final value of  $10.8 \pm 0.4\%$  in taps undergoing the CHT\_8 heat treatment. Irrespective of the type of heat treatment, there were almost no changes in mean apparent grain size, which ranged from 2.6 to 2.7  $\mu$ m (G14, according to the EN ISO 643:2020 standard). The mean plane strain fracture toughness of taps initially increased up to  $32 \pm 6$  MPa·m<sup>1/2</sup> with increasing pressure of quenching gas, but then decreased with a further increase in the same parameter, reaching a minimum value of  $26 \pm 4$  MPa·m<sup>1/2</sup>. Concerning Vickers hardness, the mean values ranged from  $892 \pm 11$  (corresponding to 66 HRC) to 906  $\pm 4$  (corresponding to 67 HRC) with varying pressure of quenching gas.

**Table 1.** Mean values of retained austenite (RA), apparent grain size, plane strain fracture toughness ( $K_C$ ) and Vickers hardness (HV1) of taps after the different heat treatments.

Parameter	Heat Treatment		
	CHT_5	CHT_6	CHT_8
RA (%)	$12.4\pm0.5$	$11.2 \pm 0.2$	$10.8\pm0.4$
Apparent grain size (µm)	$2.6 \pm 0.1$	$2.7 \pm 0.2$	$2.6 \pm 0.2$
Kc (MPa⋅m <sup>1/2</sup> )	$31 \pm 5$	$32 \pm 6$	$26 \pm 4$
HV1	$892 \pm 11$	$906 \pm 4$	$898 \pm 7$

The relationships between plane strain fracture toughness and the characteristics of small secondary carbides are shown in Figure 9. Considering the CHT\_5 and CHT\_6 heat treatments, plane strain fracture toughness increased with increasing area fraction (Figure 9a) and population density

(Figure 9b), but increased with decreasing of the size of small secondary carbides (Figure 9c). On the contrary, for taps undergoing the CHT\_8 heat treatment, the higher area fraction and population density and the smaller size of small secondary carbides resulted in the lowest plane strain fracture toughness.



**Figure 9.** Relationships between plane strain fracture toughness and (**a**) area fraction, (**b**) population density and (**c**) size of small secondary carbides.

#### 4. Discussion

The microstructure and mechanical properties of taps made of M35 high speed steel and vacuum heat treated at increasing pressures of quenching gas were studied for improving their fracture toughness and durability. The extensive metallographic investigation of taps, vacuum heat treated with pressure of quenching gas of five bar, shows a microstructure similar to that of high speed steels [2]. After tempering, the microstructure consists of a non-uniform distribution of large and dendritic-type MC and  $M_6C$  primary carbides and a fairly uniform distribution of alloyed cementite and  $M_7C_3$ secondary carbides in a tempered martensite matrix. Previous research showed that the V-rich MC primary carbides are known to be the hardest phase, main responsible for the wear resistance of HSS, whereas the W- and Mo-rich  $M_6C$  primary carbides derive from the conversion of metastable  $M_2C$  in more stable  $M_6C$  during reheating for hot-forging or hot-rolling [27–29]. Concerning the Cr-rich  $M_7C_3$ secondary carbides, studies on the effects of tempering conditions on secondary hardening of high speed steels proved that at tempering temperatures above 550 °C, these carbides gradually precipitate from the edge of the  $M_6C$  primary carbides towards the interior [30]. After this process,  $M_7C_3$  and  $M_6C$  can coexist in the matrix, as supported by our SEM/EDS and XRD examinations. Based on the results of the image analysis, the area fraction, population density and size of large secondary carbides are between 2.53 and 2.99%, 3.5 and  $4.5 \times 10^3$ /mm<sup>2</sup>, 2.66 and 2.90  $\mu$ m, respectively. Considering the characteristics of small secondary carbides, area fraction is between 2.38 and 2.68%, while population

density ranges from 130 to  $138 \times 10^3$ /mm<sup>2</sup>. The size is between 0.66 and 0.82 µm. All values of both large and small secondary carbides are comparable to those previously reported for vacuum heat treated high speed steels [18,31,32].

In the tempered martensite matrix, an average of 12.4% of retained austenite is detected, which represents good average value for commercial practice [33]. A certain amount of proeutectoid carbides is also visible along prior austenitic grain boundaries due to quench embrittlement [2]. This phenomenon involves the precipitation of carbide particles during cooling from austenitizing temperature, but when the material is still in the austenite field. The SEM/EDS analyses confirm that the precipitation occurs at the austenite grain boundaries since those regions are preferential sites for nucleation in terms of space and thermodynamic energy [2]. Overall, the presence of proeutectoid carbides does not affect the hardness of steels, but may favor crack propagation, and therefore, reduce the final fracture toughness [2]. In this study, not only the mean values of apparent grain size and Vickers hardness after tempering but also those of plane strain fracture toughness are comparable to those of high speed steels undergoing conventional vacuum heat treatments, despite the occurrence of the quench embrittlement phenomenon [9,33].

After tapping tests, failure analysis of vacuum heat treated taps by optical and scanning electron microscopy shows that, in normal operating conditions, the predominant fracture mechanism is quasi-cleavage [31,32]. Fracture initiates by cracking of primary carbides and cracking at the primary carbides/matrix interfaces, because these carbides are large and prone to fracture. Moreover, primary carbides are found in the interdendritic regions and such non-uniform distribution facilitates their cracking from the matrix at low-stress levels [20]. The nucleation of microvoids by decohesion of secondary carbides then occurs, followed by cleavage fracture. Unlike primary carbides, no cracking of secondary carbides is detected. Formation of microvoids by decohesion of secondary carbides is detected. Formation of secondary carbides when the local interfacial stresses, occurring on a carbide particle, reach a critical interfacial strength, which is a characteristic of the material.

Vacuum heat treatment of taps at pressures of quenching gas of six and eight bar to improve their fracture toughness and durability produces microstructures similar to those of taps vacuum heat treated at five bar, but with lower amounts of retained austenite and proeutectoid carbides along prior austenitic grain boundaries. An increase in the pressure of quenching gas also modifies the precipitation behavior of secondary carbides, as proved by the rise in their area fraction and population density, and by the decrease in size of large and small secondary carbides [31,32]. Slight differences are found in the mean apparent grain size and Vickers hardness of all taps.

With respect to plane strain fracture toughness, it is known that the microstructure has a great influence on this parameter [33]. In our taps, among all microstructural constituents primary carbides appear identical regardless of the type of heat treatment, since their characteristic are controlled only by the temperature and time of austenitization [32]. Accordingly, any variation of the measured plane strain fracture toughness is supposed to be due to alterations of the secondary carbides, and more specifically, the small secondary carbides. By comparing the plane strain fracture toughness of conventionally heat treated taps with that of taps vacuum heat treated at six bar, it was observed that the plane strain fracture toughness slightly increased with increasing area fraction and population density, but also with decreasing size of small secondary carbides. Based on the metallographic analyses by optical microscopy of our fracture surfaces, this may be due to the fact that small secondary carbides, more uniformly distributed, are able to deflect more effectively cracks propagation, leading to more energy absorption associated with cracking. Moreover, some authors also demonstrated that the precipitation of a higher amount of small secondary carbides reduced the contents of dissolved carbon and alloying elements in the matrix, thus raising its ductility and fracture toughness [31]. Nevertheless, taps vacuum heat treated at a pressure of quenching gas of eight bar showed the lowest plane strain fracture toughness, despite the highest area fraction and population density, and the smallest size

of small secondary carbides. This is probably due to an excessive amount of finer small secondary carbides, which may reduce the plane strain fracture toughness of our high speed steel by providing a preferential path for crack propagation.

## 5. Conclusions

This study investigates the microstructure and mechanical properties of taps made of M35 high speed steel and vacuum heat treated at increasing pressures of quenching gas, in order to improve their fracture toughness and durability. Based on the results, the following conclusions can be drawn:

- The microstructure of taps undergoing vacuum heat treatment at a pressure of quenching gas of five bar is similar to that of high speed steels. The microstructure mostly consists of primary carbides (MC and M<sub>6</sub>C), alloyed cementite and secondary carbides (M<sub>7</sub>C<sub>3</sub>) in a tempered martensite matrix, with retained austenite and proeutectoid carbides along prior austenite grain boundaries. All values of the characteristics of secondary carbides, as well as those of retained austenite, apparent grain size, Vickers hardness and plane strain fracture toughness are comparable to previously reported data for vacuum heat treated high speed steels;
- in normal operating conditions the predominant fracture mechanism of taps is quasi-cleavage. Fracture initiates by cracking of primary carbides at the primary carbides/matrix interfaces and by nucleation of microvoids by decohesion of secondary carbides, followed by cleavage fracture;
- vacuum heat treatment of taps at pressures of quenching gas of six and eight bar produces
  microstructures similar to those of taps vacuum heat treated at a pressure of quenching gas of five
  bar, but with lower amounts of retained austenite and proeutectoid carbides along prior austenitic
  grain boundaries. An increase in pressure of quenching gas also modifies the precipitation
  behavior of secondary carbides, whereas slight differences are found in the mean apparent grain
  size and Vickers hardness of all taps. In relation to plane strain fracture toughness, taps vacuum
  heat treated at six bar shows the highest values of this parameter thanks to a higher content of finer
  small secondary carbides, which deflect crack propagation. Conversely, the lowest plane strain
  fracture toughness of taps vacuum heat treated at eight bar may be due to an excessive amount of
  finer small secondary carbides, which may provide a preferential path for crack propagation.

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