Attempts have been made to describe the influence of WC additions on properties of M3/2 high speed steel (HSS) based composites. The powder compositions used to produce skeletons for further infiltration were M3/2, M3/2+10%WC and M3/2+30%WC. The powders were cold pressed at 800 MPa. The green compacts were subsequently sintered for 60 minutes at 1150°C in vacuum. These as-sintered specimens were used for copper infiltration. A qualitative EDX analysis revealed presence of both MC type vanadium-rich carbides and M₆C type tungsten and iron rich carbides. In specimens containing 10 and 30% WC the carbide phase was uniformly distributed within copper-rich regions. The WC monocarbide reacts with the surrounding HSS matrix and forms a carbide grain boundary film. The microstructural observations of the as-sintered specimens was followed by Brinell hardness test and supplemented with dilatometric studies.

Keywords: M3/2 High speed steel (HSS), tungsten carbide WC, copper, metal matrix composites (MMC), sintering, infiltration

Badania miały na celu określenie wpływu węgliku wolframu WC na własności kompozytów na osnowie stali szybkotnącej. Do wytwarzania porowatych kształtek stosowano następujące mieszanki proszków: 100% M3/2, M3/2+10%WC i M3/2+30%WC. Mieszanki proszków prasowano pod ciśnieniem 800MPa. Część wyprasek poddano spiekaniu przez 60 minut w temperaturze 1150°C w próżni lepszej od 10⁻². Do infiltracji stosowano wypraski i porowate spieki. Analiza przy użyciu mikrosondy rentgenowskiej potwierdziła obecność dwu typów węglika: WC bogatych w wolfram i żelazo. W kształtach z dodatkiem 10 i 30% węgliku wolframu następuje rozpuszczanie węgliku i rekcja z osnową stali szybkotnącej, w wyniku czego powstaje nowy węglik w postaci wilgotnego skrajku dodanego węgliku wolframu. Dodatkowo przedstawiono wyniki badań twardeści, wytrzymałości na zginanie oraz odporności na zużycie ścierne badanych kompozytów.

1. Introduction

Design criteria for high strength tool materials are: high hardness, good wear resistance and adequate toughness. Cold compaction and vacuum sintering of PM high speed steels to full density is now a well established technique [1-3]. In recent years, extensive work has been undertaken to sinter metal matrix composites (MMCs) which contain ceramic particles embedded in a HSS matrix. Most studies have been focused on sintered HSSs with additions of hard ceramics such as Al₂O₃, VC, NbC, TiC, WC and TiN with the aim of producing a more wear resistant HSS-base material [4-18]. These composites have been developed for harsh wear applications as an attractive alternative to expensive cemented carbides. They have higher hardness, elastic modulus and resistance to wear than high strength steels, however, the addition of brittle ceramic particles may degrade toughness as the reinforcing particles promote crack initiation. In order to ensure good bonding at the ceramic/matrix interfaces, the ceramic particles must exhibit some reactivity with the matrix. Except for Al₂O₃, which does not react with the HSS matrix, diffusion of iron into the MC carbide particles give rise to good cohesion across the ceramic/matrix interface. Besides, MC carbides are relatively stable in contact with iron during sintering and do not dissolve extensively. Therefore, MC particles were chosen as the reinforcement.

An inexpensive and easy route to develop HSSs reinforced with MC carbides consists of mixing commercial HSS powders with carbides. High hardness, mechanical strength, heat resistance and wear resistance of HSSs make them an attractive material for manufacture of MMCs. HSSs comprise a family of alloys mainly used for cutting tools. Their name – high speed steel – is a synthesis of the following two features [11]:

- the alloys belong to the Fe-C-X multicomponent system, where X represents a group of alloying elements in which Cr, W, Mo, V and Co are the principal ones;
- the alloys are characterized by their capacity to retain high hardness when exposed to elevated temperatures resulting from cutting metals at high speeds.

The carbides are mainly formed by strong carbide formers, i.e. V, W, Mo and Cr. Depending on the alloy composition
and its thermal processing parameters (sintering temperature and cooling rate) various types of carbides, such as $M_6C$, $M_2C$, MC and $M_23C_6$, can be formed. The carbides can grow directly from the melt during solidification or they are created by eutectic reaction or by decomposition of other types of carbide. The latter case is well exemplified by the decomposition by eutectic reaction or by decomposition of other types of carbide. For a given amount of V, increased W concentrations favour formation of $M_6C$ at the expense of MC, whereas Mo has the opposite effect. The volume fraction of eutectic carbides increases in proportion to the carbon content. Many studies demonstrate how the as-sintered microstructure can be tailored by adding elemental powders to a gas atomised $M_3/2$ HSS powder. Applications of elemental Si to repel C from the melt during sintering, elemental Ni to stabilize austenite, and elemental V to form MC at the expense of cementite are well documented [15-21]. Additions of carbides to HSSs have also been studied [9-19]. Thermodynamically less stable carbides, such as SiC and Cr$_3$C$_2$, easily dissolve in the matrix during sintering or annealing. Intermediate carbides, such as WC, VC, Mo$_2$C and NbC, react with the matrix to produce other carbide phases with compositions similar to primary carbides, e.g. $M_6C$ (Fe$_3$W$_2$C or Fe$_3$Mo$_2$C) and niobium or vanadium rich MC type carbides. Thermodynamically stable carbides such as TiC are retained in their original form but also encourage formation of MC carbides at the TiC/matrix interface. Generally TiC additions decrease sinterability by raising the sintering temperature required to achieve a pore-free material. It has been reported that additions of TiC decrease bending strength of HSSs and cause negligible increase in hardness [18,19].

### 2. Experimental procedure

Water atomised $M_3/2$ grade powder, finer than 160 $\mu$m, was delivered by POWDREX SA in as-annealed condition. Its microstructure consists of fine carbides embedded in a martensitic/bainitic matrix and particle microhardness $HV_{0.005} = 284 \pm 17$ [5].

The chemical composition of the $M_3/2$ grade powder is given in Table 1.

<table>
<thead>
<tr>
<th>C</th>
<th>Cr</th>
<th>Co</th>
<th>Mn</th>
<th>Mo</th>
<th>Ni</th>
<th>Si</th>
<th>V</th>
<th>W</th>
<th>Fe</th>
<th>O</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.23</td>
<td>4.27</td>
<td>0.39</td>
<td>0.21</td>
<td>5.12</td>
<td>0.32</td>
<td>0.18</td>
<td>3.1</td>
<td>6.22</td>
<td>balance</td>
<td>0.0626</td>
</tr>
</tbody>
</table>

Commercial tungsten carbide, having a mean particle size of 3 $\mu$m, was used as an additive. The particle shape of the experimental powders is shown in Fig. 1.

![SEM micrographs](image)

**Fig. 1.** SEM micrographs of: a) water atomised HSS M3/2 powder, b) fine WC powder

The experimental powders were used to prepare three composite materials. Their powder compositions are presented in Table 2.

### TABLE 2 Powder compositions of the experimental composites

<table>
<thead>
<tr>
<th>Material designation</th>
<th>Powder composition, wt-%</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$M_3/2$</td>
</tr>
<tr>
<td>M</td>
<td>100</td>
</tr>
<tr>
<td>M10WC</td>
<td>90</td>
</tr>
<tr>
<td>M30WC</td>
<td>70</td>
</tr>
</tbody>
</table>

The powders were mixed in a Turbula type mixer for 30 minutes and cold pressed in a rigid cylindrical die at 800 MPa. Green compacts were subsequently sintered for 60 minutes at 1150°C in vacuum better than 10$^{-2}$ Pa. The sintering process was concurrently studied by means of dilatometric measurements.

Both green and as-sintered densities of the porous bodies were measured prior to infiltration using the Archimedes method. The measured densities were divided by theoretical densities ($\rho_t$) to calculate relative densities. The theoretical densities for the composites were calculated using the following equation:

$$\rho_t = (\rho_{WC} \cdot X_a + \rho_b \cdot X_b)$$  \hspace{1cm} (1)

where $\rho_{WC}$ and $X_a$ are density and volume fraction of WC; similarly $\rho_b$ and $X_b$ are density and volume fraction of $M_3/2$ HSS.

Prior to infiltration the porous skeletons were also tested for Brinell hardness.

The infiltration process was carried out in vacuum better than 10$^{-2}$ Pa. Both green compacts and preforms, sintered for 60 minutes at 1150°C in vacuum, were infiltrated with copper. Carefully pre-weighed pieces of copper infiltrant were placed on top of the rigid skeletons of predetermined porosity, heated to 1150°C, held at temperature for 15 minutes, and cooled down with furnace to room temperature.

After infiltration the composites were examined microscopically by means of the light microscopy (LM) and scanning electron microscopy (SEM). The phase composition was examined by both the energy-dispersive X-ray spectroscopy (EDX) and X-ray diffraction (XRD).
3. Results and discussion

The combined effects of WC content and powder processing route on relative density, shrinkage and Brinell hardness of porous skeletons are shown in Figures 2, 3 and 4.

Fig. 2. Relative density of green compacts and sintered porous skeletons as a function of WC addition

Fig. 3. Volume changes of compacts during sintering as a function of WC addition

Figures 2 and 3 show the effect of WC content on compressibility and volume changes during sintering of the M3/2 HSS skeletons. The green density of compacts slightly decreases with increasing WC content, whereas additions of tungsten carbide, up to 30%, increase the as-sintered density and Brinell hardness of the HSS-base composites (Figure 4).

Fig. 4. Brinell hardness of sintered porous skeletons as a function of WC addition

Figure 5 shows that the M3/2 grade HSS cannot be fully densified at 1150°C, and that the as-sintered density is approximately equal to the green density. Additions of 30% tungsten carbide increase the as-sintered density presumably due to the occurrence of a liquid phase due to dissolution of tungsten carbide particles in the matrix. As exemplified in Figure 5, marked specimen expansion followed by its rapid contraction has indicated that the liquid appears between 1080 and 1110°C. Intermediate carbides, such as WC, partially dissolve and react with the steel constituents to produce a new carbide phase with a composition similar to primary carbides present in HSSs.

Fig. 5. Dilatometric curves recorded for the M3 and M30WC skeletons

4. Phase analysis

Phase identification of the composites was performed using a Tur 62 X-ray diffractometer and Cu Kα radiation (λ = 1.5406Å).

Fig. 6. XRD patterns recorded for as-sintered M and M30WC skeletons

From the XRD patterns in Figure 6 it is evident that M₆C (Fe₅W₃C) and MC (V₈C₇) carbides, ferrite and austenite are present in the as-sintered M sample, whereas M₆C (Fe₅W₃C and Fe₃W₃C-Fe₅W₃C) and MC (WC and V₈C₇) carbides, ferrite and austenite have been found in the M30WC sample. As compared to the M material the intensity of Feα peaks in
pre-sintered M30WC samples is smaller at the expense of WC and Fe₃W₃C-Fe₄W₂C peaks.

The SEM-EDX analysis performed on specimens containing 10 and 30% WC have revealed the carbide phase evenly distributed within the steel matrix. From Figure 7 it is evident that during sintering of porous skeletons the added WC reacts with the HSS matrix to form iron-rich Fe₃W₃C-Fe₄W₂C carbides.

As shown in Figures 8÷10, the as-infiltrated properties of the investigated composites depend on their manufacturing route and WC additions.

From Figure 8 it is evident that the molten copper is drawn into the interconnected pores of the skeleton, through a capillary action, and fills virtually the entire pore volume to yield the final density exceeding 97% of the theoretical value. Infiltration of as-sintered skeletons with copper results in slightly higher relative densities of M and M10WC composites due to reduction of oxides present at powder surface.

Fig. 7. Microstructure of the M30WC composite and EDX spectra

As shown in Figures 8÷10, the as-infiltrated properties of the investigated composites depend on their manufacturing route and WC additions.

The Brinell hardness of the as-infiltrated composites increases with the WC addition. Slight difference in hardness between the materials obtained from the two manufacturing routes was observed, i.e. higher hardness was achieved after infiltration of green compacts. This can be explained by the reduction of carbon content taking place during pre-sintering. The Bending Strength of the as-infiltrated composites decreases with increasing the content of hard WC powder in the mix.

The wear tests were carried out using the block-on-ring tester (Figure 11).

During the test a rectangular 20×4×4 mm wear sample (1) was mounted in a sample holder (4) equipped with a hemispherical insert (3) ensuring proper contact between the test sample and a steel ring (2), heat treated to 55 HRC, which was rotated at a constant speed of 500 rpm. The wear surface of the sample was perpendicular to the loading direction. Double lever system was used to force the sample towards the ring at 165 N ±1%. The loss of sample mass was measured after a sliding distance of 1000 m.

Before testing for wear, all specimens were polished to an average roughness of \( R_a = 1 \mu m \). The tests were carried out at room temperature, keeping a relative humidity below 30%. The loss of mass of the wear samples is given in Figures 12÷13.
Fig. 12. Loss of mass of as infiltrated composites

Fig. 13. Friction coefficient of as infiltrated composites

Generally direct infiltration of green compacts with copper results in higher wear resistance and lower friction coefficient of the as-infiltrated M and M30WC composites, although the friction coefficients are not highly influenced by the WC additions and the obtained wear data seems to be markedly affected by the sample preparation and experimental conditions.

5. Microstructures

Typical microstructures of copper infiltrated green compacts and pre-sintered skeletons are shown in Figures 14-16.

Fig. 14. Microstructures of M based composites, SEM

Fig. 15. Microstructures of M10WC based composites

Typical microstructures of as-sintered M3/2 baseline material are composed of a martensitic matrix, retained austenite, M₆C and MC type carbides [18,19] both located within the grains and forming a grain boundary film. As can be seen in Figures 14-16, the microstructure of the M3/2 based composites infiltrated with copper is similar except for the copper areas. Regardless of the infiltration route the tested composites have almost pore-free microstructure. The WC powder admixed to the M3/2 grade HSS markedly influenced the microstructure of the composites. Some WC particles were retained in the microstructure but a part of the tungsten monocarbide came into reaction with the M3/2 steel components to form vanadium and iron containing M₆C carbides (Figure 7). These M₆C had a tendency to form a continuous layer surrounding the added WC particles.

6. Conclusions

- Direct infiltration of green compacts with copper results in the higher hardness and wear resistance of composites and allows to cut the production cost.
- Typical microstructure of the M10WC and M30WC composites infiltrated with copper is composed of tempered martensite, residual austenite, copper regions, MC and M₆C carbides.
- The morphology and distribution of carbides depend on WC contents and production process parameters. Iron-rich M₆C type carbides (Fe₅W₃C-Fe₄W₂C) are formed during sintering as a result of chemical reaction between the added WC and matrix of the M3/2 steel.
- The mechanical properties of the tested composites strongly depend on the amount of WC added. The additions of WC increase hardness and decrease bending strength of the composites.
- The wear resistance of the tested composites is very high but the results are inconsistent and seem to be affected by the testing conditions.

Acknowledgements

This work was financed by Ministry of Science and Higher Education through the project No 11.11.110.788.

REFERENCES


Received: 20 March 2012.