Corrosion and thermal shock resistance of metal (Cu, Al) matrix composites reinforced by SiC particles

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Abstract. This paper presents the results of studies concerning the production and characterization of Al-SiC/W and Cu-SiC/W composite materials with a 30% volume fraction of reinforcing phase particles as well as the influence of corrosion and thermal shocks on the properties of selected metal matrix composites. Spark plasma sintering method (SPS) was applied for the purpose of producing these materials. In order to avoid the decomposition of SiC surface, SiC powder was coated with a thin tungsten layer using plasma vapour deposition (PVD) method. The obtained results were analysed by the effect of the corrosion and thermal shocks on materials density, hardness, bending strength, tribological and thermal properties. Qualitative X-ray analysis and observation of microstructure of sample surfaces after corrosion tests and thermal shocks were also conducted. The use of PVD technique allows us to obtain an evenly distributed layer of titanium with a constant thickness of 1.5 μm. It was found that adverse environmental conditions and increased temperature result in a change in the material behaviour in wear tests.

Key words: Metal-matrix composites, silicon carbide, wear resistance, corrosion, thermal shocks.

1. Introduction

Metal matrix composites have advantages over the conventional materials. In comparison to monolithic materials, MMCs have high strength-to-density and stiffness-to-density ratios, better fatigue resistance at room and elevated temperature [1, 2]. In addition, MMCs possess lower coefficients of thermal expansion (CTE) and better wear resistance in comparison to conventional materials. As a thermal management material, MMCs fully meet the requirement of high thermal conductivity, high thermal stability, high dimensional stability, low CTE and high processing temperature resistance [3]. Over the last few years MMCs have found many practical applications. They are used in production of engine parts, airplane fuselages, sensors, electronic packaging, etc. [4–6]. Depending on the future applications of the designed material, the key requirements are different, e.g. materials for use in the electronic industry should be characterized by the following properties: high thermal conductivity, thermal expansion coefficient adapted to the properties of semiconductors, good mechanical properties, ability to bond, stability of structure across heat cycles, and low fabrication cost [7]. Highly demanding specifications from the automotive and aerospace industry concerning fuel efficiency and emission standards on the one hand, and high standards concerning comfort and safety on the other hand, are the motivation for decreasing the weight of vehicles by using such composites. The main development objectives are an increase in stiffness, wear resistance, as well as oxidation resistance at an elevated temperature [8, 9]. Aluminium (Al) and copper (Cu) reinforced by SiC are used in various applications due to their excellent thermo-physical properties such as low coefficient of thermal expansion, high thermal conductivity, and improved mechanical properties. Cu-SiC and Al-SiC composites enable high thermal and electrical conductivity and high toughness of metal matrix to be combined with high stiffness, hardness, and wear resistance of silicon carbide reinforcement [1]. The fact that such a combination of properties is possible makes these MMCs particularly interesting for wear applications. Examples include automobile brakes, cylinder bores and bearings of internal combustion engines. Al-MMCs reinforced with silicon carbide particles are chosen because of their low cost and low specific weight compared to other types of materials. One of the most important stages for producing MMCs by powder metallurgy is the sintering process. Among the various sintering methods, spark plasma sintering (SPS) technique is one of the most effective processes in consolidating materials [7–11]. Due to the fact that the generated heat in SPS process concentrates only on particle surface, grain growth is limited. When the electrical current flows, the whole system is heated via the Joule’s effect. As a result, an increase in local temperature on grain-to-grain contact points is observed. This influences the speed of mass transport and the evaporation and convection processes. At the same time, oxides are removed from the surface of metal particles and the diffusion activation energy in the powder is decreased. This occurrence improves the densification of the material [12, 13]. It is possible to produce MMCs with planned properties by choosing the types and number

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of aluminium alloys and their reinforcement. In [14] it was reported that if unproblems appearing during the fabricating of Al-SiC composites is the formation of Al₄C₃ at the SiC/matrix interface as a result of interfacial reaction between Al and SiC (4Al + SiC → Al₄C₃ + 3Si). The Al₄C₃ phase is undesirable due to the fact that, apart from its fragility, it easily reacts with water vapor to form gaseous products that are responsible for the microcracking and, consequently, the destruction of materials (Al₄C₃ + 12H₂O → 4Al(OH)₃ + 3CH₄↑). In order to protect SiC particles before the aggressive attack of liquid Al, barrier coatings (SiO₂) are used or the amount of silicon in the matrix is increased [15]. A different situation takes place with molten Cu that decomposes SiC to Si and C at an elevated temperature [16]. According to [17], SiC decomposes in contact with Cu during the sintering process and due to this Si diffuses into the copper matrix and remaining carbon atoms form a layer. The solubility of Si in Cu is approximately 5at.% at 850°C [18]. Additionally, carbon does not react with copper and, because of the low coefficient of friction, reduced strength of the interface is observed [19]. Therefore, special attention should be paid to the phenomena in the area of metal/ceramic interface. The consolidation process should assure the maximum densification of the material, but without any structural changes in the interface area. On the other hand, there are attempts to improve the properties of Cu/SiC interface by the introduction of an additional coating on reinforcement. Coating technique has been intensively studied for A-MMCs and Cu-MMCs. In addition, various properties of ceramics and metals, especially a large difference in the values of their coefficient of thermal expansion, adversely affect the quality of the bonding of the metal matrix and the reinforcement. In such a structure, thermal stresses may occur at the interface, which may result in a decrease in the material's mechanical properties. Only a good connection enables smooth transfer of, e.g. heat flux or stress from the matrix to the reinforcing phase, and thus obtaining very good mechanical and functional properties of the composites. To achieve this, the surface of ceramic reinforcement must be protected. Recently, we have demonstrated that the use of Cr, W, Mo as coatings improves the structure and properties of metal/ceramic interfaces [16].

This work reports the results of the examination of structure and properties of Cu-SiC and Al-SiC composite materials, containing silicon carbide particles, coated with a thin tungsten layer by plasma vapour deposition technique. For application in the automotive industry, the thermal and mechanical properties of the materials were determined after their exposure to a corrosive environment and to variable heat loads. The effects of corrosion and thermal shock tests on the material density, hardness, bending strength, tribological and thermal properties were analysed.

2. Experimental procedure

In this work, copper powder produced by NewMet Koch, with a grain size of 40 µm, purity of 99.99% (Fig. 1a) and aluminium powder product by Goodfellow with a grain size < 60 µm, purity of 99.99% (Fig. 1b) were used as the matrix starting materials. Silicon carbide particles (Fig. 1c) with a mean size between 40–80 µm (Fig. 2a) (product of Saint Gobain) and purity 99.99% were selected as the reinforcement. The SEM image of SiC powder particles (Fig. 1c) distinguishes two general types of particles. The first group of particles having full 3D symmetry, i.e. DX ≈ DY ≈ DZ, and those that have 2D symmetry, type DX ~ DY >> DZ, (flattened particles). X-ray analysis of SiC powders (Fig. 2) showed that it is a material consisting of 88vol.% SiC 6H and 12vol.% SiC 15R.
To protect SiC particles against decomposition they were covered with a tungsten layer of 1.5 µm thickness using plasma vapour deposition method (PVD). In order to ensure a uniform powder coating, a special spot for powder mixing in vacuum chamber was set up. The process of spraying was conducted in ULVAC sputtering system (Figs. 3a, 3b). The process was conducted for 12 h at the power of 200 W emitted by the installation’s high-frequency generator. The morphology of the silicon carbide coated by tungsten is shown in Fig. 3c.

Powder mixture with 30vol.%SiC/W particles with aluminium and copper matrices were prepared using a planetary ball mill (Pulversette 6, Fritsch GmbH, Germany). The mixing process was conducted for 4 h at a rotational speed of 200 rpm in argon atmosphere, using a vial (250 ml) and balls (Ø = 10 mm) both made of WC/Co while 2 wt.% of stearic acid was added as a process control agent. The densification of the powder mixtures of Cu-30%SiC/W and Al-30%SiC/W was conducted using spark plasma sintering method. SPS densification process was performed in a vacuum chamber (5.0×10⁻⁵ mbar) under the following conditions: sintering temperature 950°C for Cu-MMC and 590°C for Al-MMC, heating rate – 100°C/min, holding time – 10 mins, and pressure – 20 MPa.

Corrosion resistance tests in a salt chamber (environment: 50 g/l NaCl, temperature 35°C, total time 96 hours) were performed. Every 24 hours photos of the samples were taken (Fig. 4).

The developed MMCs were subjected to resistance tests for cyclic temperature changes using the testing system presented in Fig. 5. The tests were conducted in the heating-cooling regime, using a maximum temperature of 400°C for Al-SiC/W and 600°C for Cu-SiC/W composites. Changes in
the materials structure, density, hardness, bending strength and thermal properties after 500 cycles of heating-cooling cycles were analyzed.

The tribological tests were done using high temperature tribotester from Anton Paar company. The tests were performed at two temperatures: RT and 100°C. In the test Al₂O₃ balls with 6 mm diameter and force 10 N were applied. As a protective atmosphere, argon was used. Tribological properties were tested on composite samples that previously had been subjected to corrosive and thermal shocks tests. Composite samples that were not exposed to adverse chemical and thermal factors were used as the reference material. The density of the obtained composite materials was measured by the hydrostatic method. The microstructures of the materials were examined via scanning electron microscopy (SEM) using AURIRGA CrossBeam-Workstation. The hardness (HV3) was tested with a Vickers diamond indenter using a load of 29 N with a loading time of 10 s. Each indentation was placed at least 10 diagonal lengths away from the adjacent indentation. The hardness results were averaged over 10 indentations per specimen. The bending strength was examined in a ZWICK 1446 strength machine. The samples intended for the bending strength tests had a size of 15 × 1 × 1 mm, and the head load was 10 kN. The thermal conductivity \( \lambda \) of the composites was measured using the Laser Flash Analyser LFA457/Netzsch. The \( \lambda \) measurement principle was as follows. The front side of a plane parallel solid sample is heated by a short laser pulse (Nd-YAG). The heat induced propagates through the sample and causes a temperature increase on the rear surface. The increase in temperature is measured versus time by using an infrared detector. The thermal diffusivity \( a \) and, in most cases, the specific heat \( c_p \) can be ascertained using the measured signal. If the density \( \rho \) and the specific heat are known, the thermal conductivity \( \lambda \) is determined from the relation (1):

\[
\lambda = C_p \cdot \rho \cdot a,
\]

where: \( \lambda \) – thermal conductivity in W/mK, \( \rho \) – density in g/cm³, \( c_p \) – specific heat in J/g K, \( a \) – thermal diffusivity in mm²/s [20]. In order to determine the value of heat conductivity, specific heat values were assumed on the basis of the mixing rule.

Qualitative phase analysis of the materials after corrosion tests (Cu-30SiC/W and Al-30SiC/W) was performed by means of powder X-ray diffraction (XRD) method. Polycrystalline materials in the form of bulk specimens (non-powdered) were studied. The experiments were performed using a universal Rigaku SmartLab 3 kW X-ray diffractometer equipped with a Cu X-ray tube and a 1D high-speed silicon semiconductor strip detector (D/teX Ultra 250). Powder diffraction patterns were measured in the reflection Bragg-Brentano geometry (θ/2θ scan) and the continuous scanning mode was used. Furthermore, in the case of Al-30SiC/W, parallel geometry with grazing incidence of primary beam was also applied to suppress strong substrate signals. Qualitative phase analysis was performed using a PDF4+2019 database and PDXL2 Software supplied by Rigaku.

3. Results and discussion

Figure 6 represents the microstructure of Al-30SiC/W and Cu-30SiC/W composite materials obtained by SPS technique. Microstructural analysis of sintered composites using scanning microscopy focused on the assessment of the quality of bonding between ceramic grains and the metal matrix as well as the quality of the materials, i.e. the distribution of the reinforcing phase and absence of any structural defects and porosity, which could have a significant negative influence on mechanical and thermal properties of the composites. As it can be seen in these figures, SiC particles had relatively strong bonding with aluminium and copper matrices and it is apparent that the sintering temperature and sintering time in SPS method were good.
One of the most important issues in manufacturing metal matrix composites by powder metallurgy techniques is to get a good bonding of metal particles with reinforcement and to form uniform metal/ceramic interfaces [21]. Figure 7 shows the interface between SiC particles (covered by W) with the aluminium matrix, where strong bonding without any discontinuities was observed. Particular grains are well separated from one another and no SiC aggregates can be found. The average thickness of the tungsten layer was estimated to be 1.5 mm.

Composite samples after corrosion tests were subjected to X-ray diffraction tests to analyze the surface composition. Qualitative phase analysis of the Cu-30SiC/W sample indicated dominant presence of Cu phase. Silicon carbide (6H-SiC), copper oxide Cu₂O (cuprite) and a trace amount of a less crystallized tungsten carbide (WC) phase were also observed. An experimental diffraction pattern with reference lines of matching phases from a database is presented in Fig. 8. The high
The background of experimental diffraction pattern is associated with a Cu fluorescence effect. Qualitative phase analysis of the Al-30SiC/W sample indicated the presence of aluminium (Al), silicon carbide (6H-SiC), tungsten (W), WAl12 intermetallic phase and a trace amount of an unidentified phase. The unresolved wide lines indicate the presence of an amorphous phase, which may be related to the oxidation process. The experimental diffraction pattern with reference lines of matching phases from a database is presented in Fig. 8.

The results of density measurements are presented in Table 1. The theoretical density of the composites were defined for the assumed volume contents, using the density of copper $\rho_{Cu} = 8.89 \, \text{g/cm}^3$, aluminium $\rho_{Al} = 2.71 \, \text{g/cm}^3$ and silicon carbide $\rho_{SiC} = 3.20 \, \text{g/cm}^3$.

Application of SPS technique allowed to obtain a material with a high relative density of 97–99%. The resulting composites were characterized by a negligible porosity, which is also confirmed by the observation of their microstructure. Measurements of the density of composite materials that have been subjected to corrosion and thermal shocks showed a slight change in this value. For samples after corrosion tests, a decrease in density was observed. The decrease of density after corrosion tests is attributed to the changes in the surface microstructure of the composite, microcracks formation and loss of consistency of the corroded layer with the materials. Figure 9 shows the surface of the Al-30SiC/W composite after exposure to NaCl solution. Generally, it has been found that the Al-SiC composites showed better corrosion resistance when compared with pure Al matrix, which has been confirmed in [22]. It may be attributed to the fact that SiC particles remain inert in the NaCl solution. The surface analysis of elements showed that after a corrosion test on the surface of the samples, a thin layer was formed, which included such elements as Na and Cl in both examined cases. This layer is characterized by discontinuity and numerous cracks that have affected the density of this material. Figure 9a shows a SEM micrograph of Al-30SiC/W composite where preferential localized corrosion can be seen and where corrosion starts in the interior of the sample and later expands to the surface. Figure 9a shows the paths where corrosion progresses on corroded surface of composite, here SiC particles are partially detached at pits, which are present at SiC/matrix interface. Fabrication of metal-ceramic composites sometimes leads to the formation of new components on the interface between the matrix and reinforcements, which can also influence corrosion. For example, it has been reported [23] that Al$_4$C$_3$ reaction product can be formed at the SiC particles/matrix interface during the fabrication of Al-SiC composites, which can result in a severe loss of corrosion resistance. Some investigators have concluded that the increased corrosion rate is due to the formation of Al$_4$C$_3$ at the interface [24]. Figure 9a shows that the surface of the samples underwent severe degradation,
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especially along the grain boundaries. These grain boundaries provide preferential corrosion initiation sites because of the discontinuity in the surface due to the change in structure. Corrosion progresses leading to an intense porous structure near the interface. Surface analysis of elements (EDX) confirmed that a layer consisting of Na, Cl as well as NaCl salt crystals was formed on the surface of the composite materials. The resulting NaCl crystals appeared on the surface of Cu-30SiC/W as well, as shown in Fig. 9b. Copper has long been employed in many applications because of its good mechanical properties and corrosion resistance. In some highly aggressive environments copper still suffers from serious damage. For instance, in an aerated chloride medium, the corrosion of copper occurs at a noticeable rate.

Very high concentration of Cl ions results in the formation of electrolyte layer on the Cu surface. Moreover, this layer serves as a barrier against low aggressive media; this phenomenon is called the self-protective effect [25]. NaCl crystalline grows in a homogeneous manner on differently oriented single-crystal Cu surfaces [26]. Deposition on NaCl islands of different height are found on Cu, mainly single islands are present (Fig. 9b). The corrosion behaviour of copper under chloride-containing ions involves the formation of a protective layer consisting of Cu2O formed as a result of rapid diffusion of oxygen. The Cu2O layer, which is a p-type semiconductor and thus has low electrical conductivity, is mainly responsible for the high resistance [27, 28]. After a long immersion time, the protective Cu2O layer suffered various degrees of attack and can be converted into Cu2(OH)Cl and Cu2(OH)2CO3 in the presence of chloride ions. This happens after a very long period of copper exposure to a corrosive environment.

For samples that were subjected to thermal shocks, small increase in density was observed. A long process of heating materials can contribute to the further densification of the material due to the diffusion processes. Some materials obtained by techniques such as hot pressing or hot isostatic pressure are usually subjected to an additional thermal process to improve the density of these materials. As a result of reheating the material, grain growth and individual pores elimination can be observed.

Table 1 presents the results of Vickers hardness measurements. The particulate reinforcements such as SiC are generally preferred to impart higher hardness.

The coating of reinforcements with W also leads to good quality of interface characteristics and hence contributes to improving hardness. In [29] it was observed that interface discontinuities in Cu-SiC composites (SiC grains not covered with a protective layer) affected the deterioration of mechanical properties.

When 30vol.% SiC is introduced into aluminium and copper matrices, twofold increase in hardness value was observed in relation to pure Al and Cu sinters. Moreover, higher hardness values were indicated for both composite materials that were subjected to thermal shocks. In both cases, a thin layer of metal oxides appeared on the surface of the thermally explored materials (Fig. 10). It should be mentioned that thermal shock tests were conducted in the water environment and in the air at elevated temperatures. This, for sure, facilitates the formation of surface oxides and leads to surface strengthening of the material.

Hardness measurements of composites after the corrosion test indicate a slight increase in the hardness value in relation to the starting materials. As shown above, as a result of chemical interactions, oxides of higher hardness than the matrix material are formed on the surface of the composite. At the same time, numerous microcracks are created in the structure of the surface layer, which reduces the material’s ability to transfer mechanical loads. Therefore, the effect of strengthening and weakening cancel each other out and the hardness is rather stable.

It was observed that bending strength increased with the growing hardness of the samples (Table 1). The effect of increase in the mechanical strength of composites can be related with the densification of the material. The decrease of pores and other discontinuities in the materials structure leads to improvement of bending strength.

Table 2 presents the results of the coefficient of friction and the depth of the groove formed during tribological tests for all analyzed composites. The comparison of friction coefficient changes vs. time for RT and 100°C of two groups of materials under investigation is presented in Figs. 11, 12.
Table 2

<table>
<thead>
<tr>
<th>Material type</th>
<th>Tribological test at RT</th>
<th>Tribological test at 100°C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>µ</td>
<td>groove depth [µm]</td>
</tr>
<tr>
<td>Al-SiC/W</td>
<td>0.65</td>
<td>142</td>
</tr>
<tr>
<td>after corrosion</td>
<td>0.50</td>
<td>109</td>
</tr>
<tr>
<td>after thermal shocks</td>
<td>0.70</td>
<td>93</td>
</tr>
<tr>
<td>Cu-SiC/W</td>
<td>0.50</td>
<td>64</td>
</tr>
<tr>
<td>after corrosion</td>
<td>0.55</td>
<td>57</td>
</tr>
<tr>
<td>after thermal shocks</td>
<td>0.70</td>
<td>41</td>
</tr>
</tbody>
</table>

Figure 11 shows the changes in the coefficient of friction in terms of the time under the load of 10 N for composite materials after corrosion tests. It can be seen that the temperature growth during the wear process reduces the friction coefficient in all cases analyzed. The highest values of friction coefficient were recorded for samples after thermal shock tests. These composites were characterized by the highest value of hardness and along with the increase in hardness, the resistance to movement increases, and thus the friction coefficient increases. The raise in the coefficient of friction can be attributed to the oxide layer formed during thermal shock tests in air on the surface of the samples. In this case, the volume of the groove depth was the minimum.

A more stable course of wear was observed at room temperature than at an elevated temperature. The general wear mechanism is similar for all materials. This is abrasive wear based on plastic deformation. The tribofilm formed on the surface of composite materials after corrosion tests is unstable and does not significantly affect the wear process. With such a heavy load, the counter-body is worn to create additional wear products. During wear tests in Cu-SiC/W composites, numerous microcracks were observed in the friction area.

Representative images of the groove area are presented in Figs. 12, 13. The microstructure of the groove for Al-30SiC/W composite is very inhomogeneous: gaps, chipping, and accretion are present, which cause rapid friction. For Cu-30SiC/W composites, the surface of the grooves is relatively smooth.

There was a plastic deformation/abrasion consisting in covering the ceramic phase with plastic copper. Slip lines are visible along which the counter-body travelled. Individual breakouts are well-distinguished in the structure. No visible loose wear products that are pressed into the groove surface during the process.
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4. Conclusions

In this study, the influence of corrosion and thermal shock tests on the microstructure, thermal and mechanical properties of Cu and Al based composites was investigated. Because of a high reactivity of Cu-SiC and Al-SiC systems at an elevated temperature, it seems purposeful to cover the SiC powder with a protective metallic layer. Based on the results presented, the following conclusions can be drawn. By SPS technique, composites based on copper and aluminium and containing SiC reinforcement can be successfully obtained. There was no sensible sign of microcracks or porosity in both composites and strong bonding was achieved at the metal/ceramic interfaces. Owing to PVD technique, it was possible to obtain a thin and discontinuities-free tungsten coating (1.5 μm) on SiC grains. The use of this layer prevents silicon carbide from decomposing.
ing. The increase of material hardness was obtained after thermocycling tests. The oxidation of Cu-SiC and Al-SiC composites resulted in surface chemical changes. The new oxide phase was detected, which results in the increase in hardness, flexural strength, and improved wear resistance. The obtained composites are characterized by good corrosion resistance, variable heat load as well as good thermal properties and high frictional wear resistance. The adverse environmental conditions and elevated temperature result in a decrease in the thermal conductivity of the tested composites. Therefore, they can be used as wear resistance materials in automotive and aerospace industries.

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References