Phase composition and microstructure of WC–Co alloys obtained by selective laser melting

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Received: 21 September 2017 / Accepted: 18 December 2017

Abstract. Phase composition and microstructure of initial WC, BK8 (powder alloy 92 wt.% WC–8 wt.% Co), Co powders, ball-milled powders with four different compositions (1) 25 wt.% WC–75 wt.% Co, (2) 30 wt.% BK8–70 wt.% Co, (3) 50 wt.% WC–50 wt.% Co, (4) 94 wt.% WC–6 wt.% Co, and bulk alloys obtained by selective laser melting (SLM) from as-milled powders in as-melted state and after heat treatment were investigated by scanning electron microscopy and X-ray diffraction analysis. Initial and ball-milled powders consist of WC, hexagonal α-Co and face-centered cubic β-Co. The SLM leads to the formation of major new phases W2Co3C, W4Co2C and face-centered cubic β-Co-based solid solution. During the heat treatment, there occurs partial decomposition of the face-centered cubic β-Co-based solid solution with the formation of W2C and hexagonal α-Co solid solution. The microstructure of obtained bulk samples, in general, corresponds to the observed phase composition.

Keywords: Selective laser melting / additive manufacturing / WC–Co alloys / X-ray diffraction analysis / phase composition / microstructure

1 Introduction

Selective laser melting (SLM) is one of the additive manufacturing methods belonging to a class of actively developed and advanced technologies which serve to supplement traditional manufacturing [1,2]. As well as other methods of the additive manufacturing [3] SLM can involve the use of a wide range of metallic materials and produce parts for aerospace and medical applications [4]. A high cooling rate typical of this process often ensures obtaining materials with submicron and nanocrystalline grain structure, having increased strength and wear resistance [5,6]. In fact, SLM is not limited by geometric form complexity and refractoriness of material [7]. One of the examples of work with refractory materials by the SLM method is the obtaining of WC-Co bulk alloys [8,9]. WC–Co alloys are widely used in industry as cutting tools [10,11] including tools with multilayer coatings [12,13], and also can be used as wear-resistant bulk materials [14], wear-resistant [15] and anticorrosion coatings [16].

Preliminary processing and composition of initial materials influence on the quality and properties of the WC–Co alloys obtained by SLM. Using of pre-sintering [8] and mechanical alloying [17] of initial WC-Co powders allows improving the density and surface finish of synthesized layers. In works [18,19] influence of WC/Co ratio on the cracking of bulk alloys was investigated and was shown that the powder mixture with 25 wt.% WC and 75 wt.% Co can be used for SLM to obtain materials without cracks. Another important characteristic which influences on material properties during the manufacturing WC–Co alloys is phase composition. When the alloys are obtained by sintering or SLM, phases increasing fragility and decreasing fracture resistance of the material can be formed [20,21]. Main stable phases in W-Co-C system are WC and W2C double carbides, WCo3 and W6Co7 double intermetallic compounds, and W3Co3C, W3Co2C (Me6C, W4–xCo2+xC, 0 < x < 1) and W6Co6C (Me12C) η-type ternary carbides. Cobalt exists in two modifications: α-Co (hexagonal, space group P63/mmc) and β-Co (cubic, space group Fm-3m) [22,23]. In WC–Co alloys, the transformation of WC carbide to W2C, Me6C, and Me12C carbides occurs due to decreasing carbon quantity in the alloy [24,25]. In the course of obtaining WC-Co alloys, the formation of the face-centered cubic (fcc) β-Co-based supersaturated solid solution which contains a large amount of W and Co is possible [26,27]. In work [26] alloys with high cobalt content (W and C from 0 to 20 at.%, the rest is cobalt) were obtained by quenching from the liquefied state. The lattice parameter of fcc β-Co-based solid solution increased when increasing the tungsten and carbon content in alloys from 0.3585 nm for Co90W5C5 to 0.3623 nm for Co80W10C10 composition. The formation of
W₃C, Co₃W, Co₆W₆C, and Co₃W₃C phases were observed in WC-8 wt.% Co sintered alloys obtained at different sintering temperatures (800–1600°C) [20]. Sintering at 1000°C and higher leads to the formation of the fcc solid solution β-Co(WC) of WC in fcc β-Co.

The previous works [18,19] mainly considered the influences of SLM modes on the alloy microstructure formation. This work provides a detailed consideration of the phase composition changes occurring during the process of obtaining bulk WC–Co alloys by SLM and their following heat treatment. The interconnection of phase composition and microstructure of the obtained materials is shown.

2 Experimental

The following powders were used as initial materials: WC No.1 (average particle size 50–80 nm), WC No.2 (average particle size 0.5–1 μm), BK8 alloy powder (92 wt.% WC–8 wt.% Co) (average particle size 2–5 μm), Co No.1 (PK1U type, average particle size 1–2 μm), Co No.2 (average particle size 50–80 nm). Four powder compositions were obtained via processing in a Retch PM 100 planetary ball mill at the rotational speed of 200 rpm for two hours: (1) 25 wt.% WC No.1–75 wt.% Co No.1, (2) 30 wt.% BK8–70 wt.% Co No.1, (3) 50 wt.% WC No.1–50 wt.% Co No.1, (4) 94 wt.% WC No.2–6 wt.% Co. Powder compositions processed in a ball mill were applied to the substrate as suspensions. Hard-alloy BK20 (92 wt.% WC–8 wt.% Co) and steel plates were used as substrates. The liquid having been evaporated, the thickness of the applied powder layer was estimated via an optical microscope by focusing on the powder surface and the substrate surface. The difference in scale readings of the micrometer vertical screw was the thickness of the powder layer. The average thickness of the applied layer was 40 μm. Powder layers were scanned by a laser beam with the wavelength $\lambda = 1.07 \text{ μm}$, laser spot diameter $d = 100 \text{ μm}$ with various scanning step, laser radiation power was 50 W with the scanning speed of 100 mm/s. Bulk samples were manufactured by alternating application and melting of powder material.

Microstructure and chemical composition analysis of investigated materials were performed with the TESCAN VEGA 3 LMH scanning electron microscope equipped with an adaptor for elemental analysis by energy dispersive spectroscopy. X-ray diffraction (XRD) analysis was used to determine the phase composition of the samples. XRD patterns were obtained by PANalytical Empyrean X-ray diffractometer with CoKα radiation. Phase composition analysis was performed by PANalytical High Score Plus software, software [27] and ICCD PDF-2 and COD databases [28].

3 Results and discussion

XRD patterns of initial powders are given in Figure 1. XRD patterns of powder mixtures (1)–(4) obtained in a planetary ball mill are given in Figure 2. Initial and milled powders contain WC (space group P-6m2), hexagonal α-Co (space group P6₃/mmc) and cubic β-Co (space group Fm-3m). The values of lattice parameters in the initial powders are $a = 0.251 \text{ nm}$ and $c = 0.407 \text{ nm}$ for α-Co, and $0.354 \text{ nm}$ for β-Co. To obtain bulk samples by SLM, the powders were applied to the substrates. XRD patterns of substrates (BK 20 alloy, steel) are given in Figure 3. BK20 substrate contains WC, α-Co and β-Co phases. XRD pattern of steel contains the peaks of the only α-Fe phase.
Table 1. Compositions of investigated materials.

<table>
<thead>
<tr>
<th>Composition, wt.%</th>
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<th>Composition, at.%</th>
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<tr>
<td>WC</td>
<td>W_{93.9}C_{6.1}</td>
<td>W_{50}C_{50}</td>
</tr>
<tr>
<td>BK8 (92% WC–8% Co)</td>
<td>W_{86.4}C_{5.6}Co_{8}</td>
<td>W_{43.7}C_{35.7}Co_{12.6}</td>
</tr>
<tr>
<td>BK20 (80% WC–20% Co)</td>
<td>W_{75.1}C_{4.9}Co_{29.0}</td>
<td>W_{35.3}C_{35.3}Co_{29.4}</td>
</tr>
<tr>
<td>(1) 25% WC–75% Co</td>
<td>W_{23.5}C_{1.5}Co_{75}</td>
<td>W_{8.4}C_{8.4}Co_{83.2}</td>
</tr>
<tr>
<td>(2) 30% BK8–70% Co</td>
<td>W_{25.9}C_{1.7}Co_{72.4}</td>
<td>W_{9.3}C_{9.3}Co_{81.4}</td>
</tr>
<tr>
<td>(3) 50% WC–50% Co</td>
<td>W_{46.9}C_{3.1}Co_{50.0}</td>
<td>W_{18.8}C_{18.8}Co_{62.4}</td>
</tr>
<tr>
<td>(4) 94% WC–6% Co</td>
<td>W_{88.2}C_{5.8}Co_{6.0}</td>
<td>W_{43.2}C_{15.2}Co_{60.6}</td>
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![Image](image_url)

**Fig. 4.** The microstructure of surfaces of samples obtained by SLM: a – (1) 25% WC–75% Co; b – (2) 30% BK8–70% Co; c – (3) 50% WC–50% Co; d – (4) 94% WC–6% Co.

![Image](image_url)

**Fig. 5.** XRD patterns of samples obtained by SLM: a – (1) 25% WC–75% Co; b – (2) 30% BK8–70% Co; c – (3) 50% WC–50% Co; d – (4) 94% WC–6% Co.

![Image](image_url)

**Fig. 6.** XRD patterns of sample (1) 25% WC–75% Co obtained by SLM after annealing: a – 1 h at 1300°C in air; b – 1 h at 1200°C in argon.

(space group Im-3m). Table 1 demonstrates compositions of all initial materials, substrates and studied compositions in at.% and wt.%. Microstructures of surfaces of bulk samples obtained by SLM are given in Figure 4. Scanning step for samples (1) and (3) was 50 μm, for samples (2) and (4) was 100 μm; as a result, tracks with the average width of 40–50 μm (Fig. 4a and c) and 100 μm (Fig. 4b and d) accordingly were formed.

The XRD patterns of surfaces of the samples obtained by SLM are given in Figure 5. The samples (1) and (2) are a practically single phase and contain the only phase – solid solution of tungsten and carbon in fcc β-Co marked as β-Co (W,C) (Fig. 5a and b). The formation of the solid solution is testified by extended lattice period as compared to the initial β-Co [26]. β-Co(W,C) lattice parameters in the samples (1) and (2) are 0.360 and 0.361 nm accordingly. XRD pattern of the sample (1) has a peak at 2θ = 49 deg. This peak can most probably be referred to W_{4}Co_{3}C carbide phase. XRD pattern of the sample (2) contains, in addition to the peaks of the main phase, the weak peaks of which can be referred to CoWO_{4} oxide. WC phase has not been detected in the samples, which testifies its complete dissolution in β-Co as in the sample (2) and partial transformation to W_{4}Co_{3}C carbide as in the sample (1). The sample (1) was heat treated in air (1300°C, 1 h, cooling with furnace) and in argon (1200°C, 1 h, cooling with furnace). XRD patterns of the sample (1) after heat treatment are given in Figure 6. During the heat treatment in air the sample’s surface oxidizes strongly and consists mainly of CoWO_{4} and CoO oxide phases, also containing a small amount of β-Co(W,C) with the lattice parameter of 0.356 nm (Fig. 6a). After heat treatment in argon sample.
Co powder. The difference in lattice parameters of the changed lattice parameters as compared to the initial amount of W and C dissolved in (W,C) phase in the samples (1) and (2) is attributed to a formation of the (W,C) lattice parameter after heat treatment testifies partial decomposition of the solid solution. It is difficult to say anything of the decomposition because of strong oxidation of the sample (1) surface after heat treatment in air. After heat treatment in argon, α-Co and Co3W phases are formed as a result of the β-Co(W,C) decomposition. In general, the formation of the solid solution based on β-Co high-temperature phase can be explained by stabilization of fcc β-Co high-temperature modification [22] during the dissolution of tungsten carbide in cobalt; moreover, the cooling process during SLM is of quenching nature. Sample compositions and β-Co(W,C) lattice parameter in the samples (1) and (2) match the results of work [26] quite well, where fcc β-Co(W,C) supersaturated solid solution was formed in Co–W–C alloys with a high Co content. The sample (3) contains W5Co2C and β-Co(W,C) (Fig. 5c).

The lattice parameter of the β-Co(W,C) is 0.362 nm, this value corresponds to the composition of $W_{10}C_{10}Co_{80}$ [26]. The chemical composition of the sample (3) is $W_{18.8}C_{18.8}Co_{62.4}$ (Table 1). The WC tungsten carbide not found in the sample partially transformed to W5Co2C carbide and was partially dissolved in β-Co. The sample (4) contains WC and W2C phases, and CoO oxide (Fig. 5d). Traces of CoO oxides in this sample and CoWO4 in other samples (except sample (1) after heat treatment in air) is probably related to surface oxidation during the laser treatment. However, the results of chemical analysis of the sample (4) cross-section also revealed the presence of cobalt at some depth from the surface. The absence of pure cobalt peak points on the XRD pattern of the sample (4) is related to low cobalt content in the sample and low X-ray penetrability. The formation of W7Co2C, W5Co2C, and W2C carbide phases testifies the decreased carbon amount in the samples, which may be connected with its burning-out in the course of materials melting during the SLM laser treatment [22–24]. The peak points in XRD patterns which failed to be identified are marked by "?".

Microstructures of the samples (1)–(4) are given in Figure 7. Microstructures of the samples (1) and (2) do not contain any visible inclusions, which conforms to the information about phase composition and confirms practically full WC dissolution in cobalt (Fig. 7a and b). The sample (1) has a dendritic structure. Diagonal dark and light lines alternating at the distance of 50 μm can be observed on the cross-section of the sample (3). Light lines correspond to β-Co(W,C) phase. Dark lines contain a lot of submicron inclusions of W4Co2C phase which were released as a result of repeated laser heating during the parallel scanning of the next track in the powder layer (Fig. 8). The distance of 50 μm between the lines of the outlined carbide phase corresponds to the scanning step of 50 μm. The sample (4) (Fig. 7d) contains equally distributed over the whole volume micron and submicron fractions of WC and W2C carbide phases in the cobalt matrix.

4 Conclusions

This work has studied the microstructure and phase composition of WC-Co alloys obtained by SLM. To obtain bulk alloys, initial powder mixtures containing 25, 27.6, 50, and 94 wt.% WC as well as 75, 72.4, 50, and 6 wt.% Co accordingly were used. During the SLM in alloys with 75
and 72.4 wt.% Co, WC phase dissolves completely with the formation of the face-centered cubic β-Co-based solid solution β-Co(W,C) containing W and C. A small amount of W₄Co₂C (Me₆C) carbide is formed in the alloy with 75 wt.% Co. WC carbide is completely dissolved in the alloy with 50 wt.% Co transforming to W₂Co₂C (Me₆C) carbide and forming β-Co(W,C) phase. WC carbide is retained in alloy with 6 wt.% Co, and W₃C carbide is formed. The β-Co lattice parameter in initial powder and lattice parameter of β-Co(W,C) in alloys with 75, 72.4, and 50 wt.% Co are 0.354, 0.360, 0.361, and 0.362 nm, accordingly. During annealing of the alloy with 75 wt.% Co, β-Co(W,C) solid solution partly decomposes forming Co₃W phase and hexagonal α-Co-based solid solution. The formation of W₄Co₂C, W₃Co₃C, and W₂C carbides occurs due to decreasing carbon content in alloys. No inclusions are observed in the microstructure of alloys with 75 and 72.4 wt.% Co. Co₃W₁₋ₓCₓ, WC, and W₂C carbide phases in the cobalt matrix are seen in the microstructure of alloys with 50 and 6 wt.% Co. The obtained results can be used in the metalworking industry and materials science for production of hard alloys and products from it both by traditional sintering and by additive manufacturing methods.

Acknowledgements. This work was financially supported by the Ministry of Education and Science of Russian Federation in the framework of the state No.11.1267.2017/4.6. The work is carried out on the equipment of the Center of collective use of MSTU “STANKIN”.

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Cite this article as: R.S. Khmyrov, A.P. Shevchukov, A.V. Gusarov, T.V. Tarasova, Phase composition and microstructure of WC–Co alloys obtained by selective laser melting, Mechanics & Industry 18, 714 (2017)