The effects of Co addition on the mechanical and microstructural properties of the matrix material of a Fe-based marble cutting tool

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Abstract

In this study, four different kinds of marble cutting matrices with a base composition of Fe-Cu-Sn and with variable amounts of Co powder were produced. For compaction, hot sintering process was carried out at 850°C under 35 MPa for 4 min. The mechanical and microstructural properties of the matrix materials were determined using SEM with EDS, three-point bending test, density and hardness measurement techniques. The addition of Co powders to the matrix and the subsequent hot sintering process affected the mechanical and microstructural properties of the cutting tool, such as hardness, total elongation, % porosity and phase distribution.

Key words: marble cutting tools, cutting matrix material, hot sintering process, mechanical and microstructural properties

1. Introduction

Diamond cutting tools manufactured using powder metallurgy are used in a wide variety of applications such as machining of products made out of natural stone, glass, ceramic and metal, highway construction, and boring works. The demand for diamond cutting tools increases day by day. 60 % of these tools and almost all of the marble cutting tools (MCT) are manufactured using the powder metallurgy (PM) technique [1]. In this technique, production of the cutting tool is achieved by mixing diamond particles with metal powders and then compacting by variety of techniques, such as: pressureless sintering [2–6], hot isostatic pressing [7], combination of pressureless sintering and hot isostatic sintering [8], extrusion [9], laser melting [10], combination of hot compaction and laser cutting [11, 12] or another specialized technique [13]. The primary factors affecting the cost of cutting tool production are the optimum composition of the materials and selection of other processing parameters. In the production of stonecutters or drilling tools, the manufactured part is composed of approximately 20 % synthetic diamond and 80 % powders made up of various combinations of Cu, Sn, Fe, Ti, Co, Ni, Al, Mo, CuSn and SiC [14].

Marble cutting or abrasion is mainly performed by the diamond particles in a cutter. Normally, small-sized diamond particles are preferred for the cutting of the hard marbles, e.g. granite and cherry marble, while coarse-sized diamond particles are used for soft marbles. Ideally, the diamond particles should reside in their positions in the metal matrix until they are abraded. In other words, diamond particles and the metal matrix holding these particles should abrade at the same rate. If the matrix abrades later than diamond particles, the diamond particles make poor contact with the marble. In this situation, cutting or abrasion does not occur and the surface of the cutting tool heats up and burns. In the case of rapid abrasion of the matrix material, diamond particles separate from their positions and fall without cutting. For this reason, together with the cutting parameters and properties of the stone to be cut, full control of the properties of the matrix material and the cutting tool is very important [15–17].

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Table 1. Chemical composition of the specimens

| Specimens | Elements (wt.%) |
|-----------|----------------|
|           | Fe  | Cu  | Sn  | Co  |
| S1        | 78  | 20  | 2   | –   |
| S2        | 68  | 20  | 2   | 10  |
| S3        | 63  | 20  | 2   | 15  |
| S4        | 58  | 20  | 2   | 20  |

2. Experimental study

2.1. Fabrication of cutting matrix material

Hot sintering produced matrix materials with different Co content for use in experimental studies. Table 1 presents chemical composition of the materials. In the production of specimens, Fe, Cu, Co and Sn powders with 99.9 % purity and particle sizes between 1.1 and 200 µm were used. Co-free reference sample S1 produced using Fe, Cu, and Sn, was produced before the other matrix materials (S2, S3, S4).

For production, as-received raw powders were weighed by a balance with 10⁻⁵ g sensitivity in the proportions given in Table 1. The uniform mixing of these powders was achieved by using a 2 kg capacity tubular mixer having two closed chambers and capable of shaking in three dimensions. The first stage of the mixing involved mixing of powders in suitable containers for 15 min at a rate of 20 rpm. In the second stage, Cr-coated steel balls (8 mm in diameter) and 1 % polyethylene glycol (PEG) were added to the containers and samples were mixed for a further 30 min at 20 rpm.

After the mixing stage, powder mixtures were poured into molds made out of graphite plates, the tops of which were covered by top compaction graphite. Graphite molds were put into a steel container and tightened smoothly with compression pins. To control the variation in temperature, the thermocouple was mounted on the mold such that the tip of the thermocouple was positioned at the center of the mold. The whole set-up was then subjected to hot compaction under a protective atmosphere. Starting from 5 MPa, samples were hot compacted at constant temperature and pressure (35 MPa at 850°C) for four minutes. Then, the heating unit was turned off, and samples were allowed to cool to room temperature under 35 MPa pressure.

2.2. Characterization of the cutting matrix material

The fabricated cutting matrix materials were characterized by density, hardness, three-point bending test and microstructural properties. The density of the matrix materials was measured according to ASTM B311-92 standard, which relies on Archimedes’ principle. The Brinell hardness of the samples was measured by using balls with 2.5 mm diameter and 62.5 kg of the load; the average of eight measurements was taken as the hardness value. The tensile strength values of the cutting tool matrix materials produced in this study were not measured by standard tensile testing specimens as the samples were much smaller than what is described in the standard. Instead, a three-point bending test was applied to find the strength values. Bending strength values of the samples were measured by a universal test machine, which is suitable to ASTM B 528-83 standard, and TRAPEZIUM X software [17].

For microstructural examination, the surfaces of the samples were smoothed with emery paper, polished and cleaned with pure alcohol. Polished surfaces were etched by an etchant mixture (HNO₃ with alcohol) so as to create visual differences between different phases. Finally, microstructural examinations were conducted by using an FE-SEM with EDS analyzer (JEOL JSM 7001F).

3. Results and discussion

Mechanical and microstructural properties of the matrix materials were determined using different techniques including SEM, EDS analysis, three-point bending test, density and hardness measurement techniques.

3.1. Density

The density measurement of samples with different amounts of Co was performed by using Archimedes’ principle. Then, the experimental values were compared with the theoretical values (Table 2 and Fig. 1).
Table 2. The theoretical and measured density values of cutting matrix material samples

| Specimens | Measured density (g cm\(^{-3}\)) | Theoretical density (g cm\(^{-3}\)) | Density (%) |
|-----------|----------------------------------|-------------------------------------|-------------|
| S1        | 7.57                             | 8.05                                | 94.09       |
| S2        | 7.85                             | 8.14                                | 96.45       |
| S3        | 7.98                             | 8.19                                | 97.42       |
| S4        | 8.00                             | 8.24                                | 97.07       |

Fig. 2. The variation of hardness value with Co content.

As depicted in Fig. 1, the measured densities of the as-sintered samples are quite close to the theoretical values. The coherence between the density values can be considered as an indication of the effectiveness of the sintering process. The difference between theoretical and experimental values of the density is an indication of the level of the porosity in the samples. According to this, samples having the least amount of porosity were S3 and S4, with Co content 15–20 %. On the other hand, sample S1 (Co-free) had the highest level of porosity among all samples examined.

Fig. 3. Results of the three-point bending test for the specimens examined in the study.

3.2. The results of hardness and bending tests

The hardness values of the samples are shown in Fig. 2. It can be clearly seen that the hardness values of the samples are increased linearly with the Co content. For example, while the hardness value of the Co-free sample S1 was measured as 120 HB, it was found as 145 HB for sample S4 (20 % Co).

The stress-strain curves of samples obtained from three-point bending test are shown in Fig. 3. The Co-free sample S1 possessed nearly 45 % elongation with a bending strength value of 850 MPa. Meanwhile, if the stress-strain graph of Co-free sample S1 is compared with that of Co-containing samples S2, S3,
and S4, it can be clearly seen that with the addition of Co, percentual elongation decreased, while bending strength increased. Based on these results, it can be stated that the addition of Co and the sintering process led to a good level of homogeneity and an optimum value of strength. Apart from this, it is already known that the presence of Cu and Sn in the matrix is responsible for the high ductility and strength of these systems [18].

### 3.3. Microstructure of the alloys

SEM pictures of samples are given in Figs. 4, 5, 6 and 7, respectively. These figures also include the results of the EDS analyses for the numbered points designated in the pictures. The dominant phases for these points are also mentioned.

According to EDS analyses of the points indicated on the picture of sample S1, the microstructure was mainly composed of $\alpha$Fe + $\beta$Cu. According to the literature, dark regions are $\alpha$Fe, and bright regions are $\alpha$Fe + $\beta$Cu [19]. In Fig. 5, which shows the SEM picture of sample S2, it can be seen that Co-containing regions appeared with the addition of Co. If the SEM pictures of samples S2 and S1 are compared, it is apparent that the sample S2 has less amount of

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**Table 3. Semi-quantitative EDS results for the label 1-3 (S1), in Fig. 4**

| Element | Weight (%) | Atomic (%) | Phase       |
|---------|------------|------------|-------------|
| Label 1 |            |            |             |
| C K     | 1.96       | 8.85       | $\alpha$Fe + $\beta$Cu |
| Fe K    | 72.56      | 70.27      |             |
| Cu K    | 23.43      | 19.95      |             |
| Sn L    | 2.05       | 0.93       |             |
| Total   | 100.00     |            |             |
| Label 2 |            |            | CuSn + Fe   |
| C K     | 2.27       | 11.23      |             |
| Fe K    | 5.58       | 5.93       |             |
| Cu K    | 84.67      | 79.10      |             |
| Sn L    | 7.48       | 3.74       |             |
| Total   | 100.00     |            |             |
| Label 3 |            |            | $\alpha$Fe + Cu |
| C K     | 1.80       | 7.86       |             |
| Fe K    | 97.34      | 91.43      |             |
| Cu K    | 0.86       | 0.71       |             |
| Sn L    | 1.87       | 0.85       |             |
| Total   | 100.00     |            |             |

**Table 4. Semi-quantitative EDS results for the label 1-3 (S2), in Fig. 5**

| Element | Weight (%) | Atomic (%) | Phase       |
|---------|------------|------------|-------------|
| Label 1 |            |            |             |
| C K     | 2.13       | 9.58       | $\alpha$Fe + CuSnCo |
| Fe K    | 62.74      | 60.58      |             |
| Co K    | 11.36      | 10.39      |             |
| Cu K    | 21.91      | 18.59      |             |
| Sn L    | 1.87       | 0.85       |             |
| Total   | 100.00     |            |             |
| Label 2 |            |            | CuSnCo + Fe |
| Fe K    | 1.92       | 2.21       |             |
| Co K    | 4.22       | 4.62       |             |
| Cu K    | 89.45      | 90.77      |             |
| Sn L    | 4.41       | 2.40       |             |
| Total   | 100.00     |            |             |
| Label 3 |            |            | $\alpha$Fe + Cu |
| Fe K    | 99.43      | 99.50      |             |
| Cu K    | 0.57       | 0.50       |             |
| Sn L    | 1.87       | 0.85       |             |
| Total   | 100.00     |            |             |
Table 5. Semi-quantitative EDS results for the label 1–3 (S3), in Fig. 6

| Element | Weight (%) | Atomic (%) | Phase          |
|---------|------------|------------|----------------|
| C K     | 2.04       | 9.16       | αCo, αFe + Cu  |
| Fe K    | 62.79      | 60.55      |                |
| Co K    | 18.00      | 16.45      | αCo, CuSn      |
| Cu K    | 15.37      | 13.03      |                |
| Sn L    | 1.80       | 0.82       |                |
| Total   | 100.00     |            |                |

Table 6. Semi-quantitative EDS results for the label 1–3 (S4), in Fig. 7

| Element | Weight (%) | Atomic (%) | Phase          |
|---------|------------|------------|----------------|
| C K     | 1.92       | 8.73       | αCo, αFe + Cu  |
| Fe K    | 55.32      | 53.99      | αFe, αCo, CuSn |
| Co K    | 21.24      | 19.64      |                |
| Cu K    | 19.46      | 16.69      |                |
| Sn L    | 2.05       | 0.94       |                |
| Total   | 100.00     |            |                |

It is apparent from the SEM pictures and EDS analyses that the two-stage mixing operation performed before the sintering process was sufficient for the homogeneous distribution of the metal powders. It is considered that the sintering process performed at 850°C is sufficient for homogeneous mixing of Co and Cu powders in solid state. It shows that the selected sintering temperature and duration were sufficient for the process. Phase separation between the 2 % Sn present and the rest of the matrix did not occur during sintering, although the melting point of Sn is about 232°C. The reason behind this is that according to the Co-Sn equilibrium phase diagram, a stable phase forms between αCo and Sn at 500–1121°C [19]. Taken altogether, SEM and EDS analyses showed that homogeneous matrix structures were obtained in marble cutting tool samples made out of Fe-Cu-Co-Sn quaternary compounds.

4. Conclusions

A marble cutting-tool matrix material made out of Fe, Cu and Sn was produced with the addition of 10, 15 or 20 % Co and using hot sintering process. Together with the microstructural examination conducted by SEM and EDS, hardness, density values, and three-point bending properties of the samples were determined. The following conclusions have been reached:

1. With the addition of Co to the cutting tool material Fe-Cu-Sn powder mixture, the density value of the tool material increased. The Co-free sample S1 had the lowest density value among all samples examined, while samples S3 and S4 with Co contents of 15 and 20 %, respectively, had the highest density values.

2. Co addition increased the hardness value of the alloy.

3. The addition of Co to the material also increased the tensile strength while decreasing the % elongation.

4. SEM and EDS analyses show that hot compaction sintering performed at 850°C for 4 min. was sufficient to obtain a homogeneous distribution of phases in the microstructure. For these reasons, this alloy, which was produced from metal powders, has sufficient qualifications to be proposed as a marble cutting-tool matrix material.

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