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Sintered Materials Based on Copper and Alumina Powders Synthesized by a Novel Method

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1. Introduction

The intensification in research of nanostructure materials that occurred in recent years was mainly due to their striking potential, i.e. mechanical and physical properties significantly improved compared to the conventional grain materials (Moriarty, 2001; Ristić, 2003).

Nano-structured materials rank in the group of ultra fine, metastable structures containing a high concentration of defects (point defects, dislocations) and boundaries (grain boundaries, interphase boundaries, *etc.*). These materials are structurally different from crystals and amorphous forms because of the fact that grain boundaries and interphases represent a specific state of the solid matter, since the atoms on boundaries are subjected to a periodical potential field of the crystal from both sides of the boundary (Koch, 1999).

The synthesis of powders represents the starting and crucial stage in the production of sintered metal materials with required properties. Considering that the starting structure undergoes certain changes during further processing, but remains essentially preserved in the structure of the final product (Ristić, 2003; Motta et al. 2004), there is an increased necessity for a great number of methods for producing powders.

Copper is widely exploited in industry because of its high electrical and thermal conductivity, even though it possesses low strength especially at elevated temperatures. In order to overcome this problem it can be strengthened by using finely dispersed particles of stable oxides like alumina, titania, yttria etc. Copper-based composite materials are widely applied in the field of electronics and electrical engineering as highly conductive materials for operation at elevated temperatures, as electrodes for resistance spot and seam welding, different contact materials, various switches, thermal and electric conductors, microwave tubes, etc (Lee & Kim, 2004).

Introduction of fine dispersed particles into matrix of base-metal has considerable strengthening effect and nano particles of oxides are especially suitable. Due to their hardness, stability at elevated temperatures and insolubility in copper they represent obstacles for dislocation, grain and sub-grain boundary movement increasing mechanical properties of these materials with insignificant effect on thermal and electrical conductivity (Naser et al., 1997; Trojanova et al., 1999, Tian et al. 2006). Significant reinforcing effects can

be kept even at elevated temperatures. For such reinforcement nano-particles of oxides are suitable (Karwan-Baczewska, 2005).

A very important aspect of dispersion strenghtening is introduction of low volume fraction of dispersed oxide particles into the volume of the base-metal, a uniform distribution of oxide particles and their fine dispersion especially in nanometer scale (Ristic & Milosevic, 2006).

Alumina was known for its exceptional properties such as high melting point, high hardness, excellent thermal stability and chemical inertness. Also, addition of alumina particles can increase the temperature of recrystallization by pinning grain and sub-grain boundaries of copper matrix and blocking the movement of dislocations highly improving strength at elevated temperatures (Plascencia & Utigard, 2005; Lianga et al., 2004). The usual amount of alumina used for dispersion strengthening is from 0.5-5.0 wt.% (Jena et al., 2004), but significant results regarding particle size could be achieved with even higher amounts ranging up to 50 wt.% alumina (Brocchi et al., 2004).

Because these oxides are not soluble in copper P/M techniques must be used instead of conventional ingot metallurgy. Oxide dispersion strengthened (ODS) copper can be successfully synthesized by following methods: highly energetic reactive milling (Ahn et al., 1996), precipitation from solution (sol-gel (Ruys & Mai, 1999), hydrothermal synthesis (Byrappaa & Adschirib, 2007), electrochemical synthesis (Yuana et al., 2007)), internal oxidation (Afshar & Simichi, 2008) etc. The production of powders by the thermo-chemical method is not a new procedure, but in recent years, due to the development of contemporary materials with pre-set properties, there has been an extended interest in this method, especially for the production of nanostructured powders (Wu et al., 2001; Jena et al. 2001; Lee et al, 2001).

Route for synthesis of composite powders based on copper and alumina presented in this research, may be regarded as new for materials in this system, even though phases of this process have been previously investigated by the authors (Korać et al., 2005; Anđić et al., 2006; Anđić et al., 2007a; Anđić, 2007b; Korać et al., 2008a, Korać et al., 2008b) and also previous attempts have been made for application of similar process in system Cu-Ag-Al₂O₃, when a three-component system was produced by mechanically alloying the thermochemically synthesized Cu-Al₂O₃ and Cu-Ag powder (Anđić et al., 2006).

Objective of this research was to investigate the possibility of copper strengthening with fine dispersed alumina particles by application of new synthesis method and production of sintered materials with properties suitable for material application as contact material.

After sintering compacts were thermo-mechanically treated in order to simulate real manufacturing process.

Verification of obtained results is performed by tribological investigations.

2. Experimental

Generally synthesis process included two separate routes. The first included synthesis of $\text{Cu-Al}_2\text{O}_3$ with 50 wt. % of Al_2O_3 (from now on $\text{Cu-50Al}_2\text{O}_3$) by thermo-chemical route. The second part of the synthesis process was mechanical alloying of atomized copper powder with powder of $\text{Cu-Al}_2\text{O}_3$ previously synthesized.

Starting raw materials for powder synthesis by thermo-chemical route were soluble salts, nitrates of copper and aluminum of p.a. quality, dissolved in distilled water (50wt.% solution) in suitable ratio for final powder to contain 50wt.% of Al_2O_3 in structure.

Copper powder used in mechanical alloying stage was produced by atomization with particle size 95% <45µm. In previous research (Korać, 2005) electrolytic copper was used for

mechanical alloying. Electrolytic copper was replaced with atomized, due to the fact that its dendrite structure prevents formation of homogenous structure of final material, even after prolonged milling time.

Stages of synthesis process included the following processes:

- 1. Spray drying of nitrate solution using Spray Dryer Büchi B-290 Advanced (Okuyama & Lenggoro, 2003) with inlet/outlet temperature 190/143°C and solution flow rate was 10% of pump power,
- 2. Dried nitrates were subjected to heat treatment in a laboratory electroresistant tube furnace (Korać et al., 2009) in air at 900°C for 1h in order to form oxides of copper and aluminium,
- 3. Oxides were then reduced in the same furnace in hydrogen atmosphere (flow rate 20L/h at $350^{\circ}C$ for 1h) in order to obtain metallic copper. Previously formed stable Al_2O_3 remained unchanged during reduction (Korać et al., 2009),
- 4. Mechanical alloying was the next step in the synthesis process. Atomized copper powder was mechanically alloyed with produced composite powder with 50wt.% Al_2O_3 in a ceramic ball mill, with dimensions ø180×160mm. Milling media were corundum balls (\geq 99% Al_2O_3 with 30mm in diameter) in order to prevent contamination of material; ball to powder ratio was 1:30. Optimal milling time was fixed at 5h for mill rotation of 300min⁻¹. Quantities of added copper powder were adjusted so that the final amount of alumina Al_2O_3 in composite powder would contain 1, 1.5 and 2 wt. %.

Flow-sheet of applied synthesis process is presented in Fig. 1.

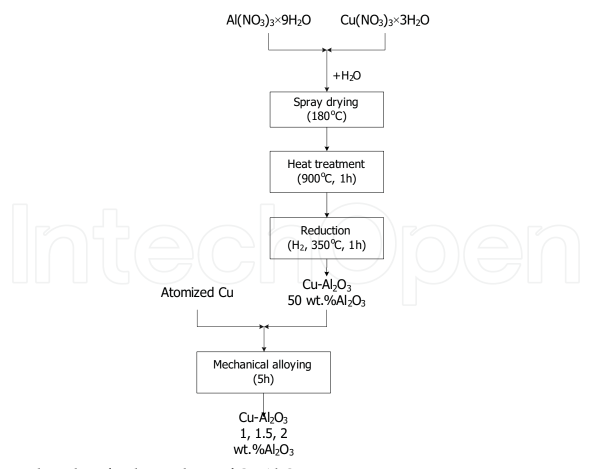


Fig. 1. Flow-sheet for the synthesis of Cu-Al₂O₃ composites

Solution concentration of 50% enabled production of higher amounts of fine particles. If solution concentration was increased coarse particles would be produced, which could disable fine coating of copper particles in later stage of mechanical alloying. On the other hand if solution was too diluted, finer particles would be obtained, but disadvantage is that in this case synthesis process would last longer, with higher equipment engagement, i.e. lower possibilities for industrial application.

Method of choice for acquiring of precursor powder was spray drying. In recent years this method is engaged in large share for production of micron and sub-micron powders (McCandlish et al., 1994). Advantages of this drying method are control of size, morphology and particle composition, as well as significant possibility of industrial application from aspect of price-productivity ratio (Okuyama & Lenggoro, 2003; Iskandar et al., 2003). Also, application of this drying method in Cu-Al₂O₃ system showed very good results in previous research (Anđić, 2006; Korać; 2007).

Temperatures of heat treatment and reduction were previously optimized for this system (Anđić, 2006), as well as milling condition (Korać, 2005).

After mechanical alloying obtained powders were compacted by a uniaxial pressing in order to minimize density gradient trough sample. Used tool had dimensions 8×32mm and height 3mm, and applied compacting pressure was 500 kN.

Sintering of samples was performed in hydrogen atmosphere in isothermal conditions at five different temperatures in the range from 725-925°C for 15 to 120 min.

After consolidation of the obtained powders, the compacted samples were uniaxially compressed by cold rolling, reduction degree of 15 and 30%. In order to determine the stability at higher temperatures, the rolled samples were annealed at temperature of 800°C for one hour in the hydrogen atmosphere.

Produced powders were characterized by X-ray diffraction analysis (XRD). XRD was performed using APD 2000-Ital structures with $CuK\alpha$ radiation, 2θ =0-100°.

Characterization of sintered samples (referred as composite in the following text) included Scanning Electron Microscopy (SEM), Energy Dispersive Spectrometry (EDS), HRF hardness measurements and electrical conductivity.

SEM analysis was performed on JEOL JSM-T20 on polished samples subsequently etched with 40 vol.% HNO₃ solution. EDS analysis was conducted on unpolished composites using Oxford system attached to JEOL SEM JSM-5800.

Ames Portable Hardness Tester was employed for hardness measurements using 1/16" ball with applied load of 60kg. For electrical conductivity measurement Foerster Sigma Test 2.069 operating at 120 kHz and with 8mm electrode diameter, was used.

Values of hardness and electrical conductivity represent the mean value of at least six measurements conducted on the same composite.

Wear testing was performed by the method of Taber in accordance with Guide to friction, wear and erosion testing (ASTM standard MNL 56, 2007). Basic information on test conditions are:

- diameter of the grinding plate 640mm,
- diameter of the grinding path 265mm,
- calculated speed rate of the grinding path 832.5mm
- granulation of Corundum Al₂O₃ Ø100μm.

3. Results & discussion

It was expected that by the proposed synthesis method nano-composite $\text{Cu-Al}_2\text{O}_3$ produced by thermo-chemical route would form homogenous compact layer around atomized copper

powder particles during mechanical alloying process. This way a ductile, highly conductive core with high strength shell layer of composite, will be achieved. In later stages this microstructure would provide dislocation blocking and prevent grain boundary motion as well as an increase of recrystallization temperature.

3.1 Powder characterization

SEM analysis of $\text{Cu-}50\text{Al}_2\text{O}_3$ presented in Figure 2 shows that powders are flake like with average particle size of 200-400nm, which have partially agglomerated due to their small size and high surface energy. Average size of present agglomerates is 1-5 μ m.

Previous results on thermo-chemical synthesis of Cu-Al₂O₃ powders (Korać, 2005) indicates that presented particles in fact represent very fine agglomerates, and that actual size of synthesized powder is in range 20-50nm.

X-ray diffraction pattern of powder containing 50 wt.% Al₂O₃ is presented in Figure 3. Identified peaks correspond to Cu and CuAl₂O₄. CuAl₂O₄ phase represents the metastable equilibria, developed in microstructure during powder synthesis process, heat treatment and reduction.

Thermo-chemically synthesized powder was used for mechanical alloying of electrolytic copper, SEM micrograph presented in Figure 4.

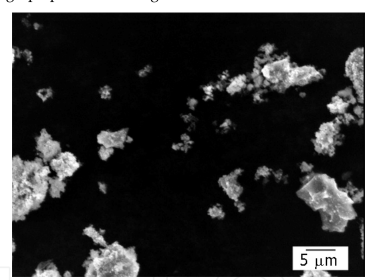


Fig. 2. SEM micrograph of Cu-50Al₂O₃ powder



Fig. 3. X-ray diffraction of Cu-50Al₂O₃ powder

Used copper powder particles are mostly spherical shaped with smooth surface. This particle morphology enables achievement of desired structure with highly conductive core with homogenous shell layer around it.

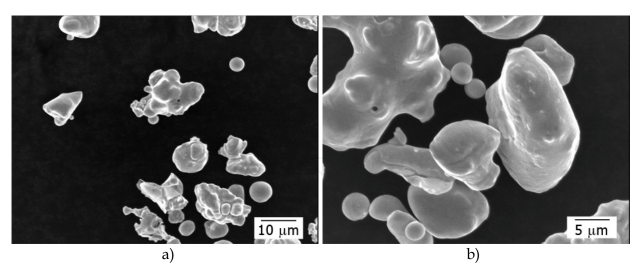


Fig. 4. SEM micrographs of electrolytic copper powder

After 5h of milling of powders, all three compositions showed structure characteristic for mechanical alloying (Figures 5-7).

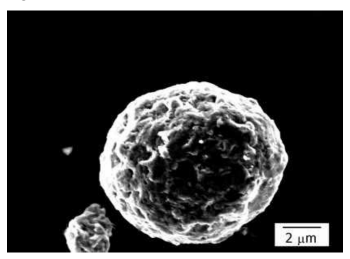


Fig. 5. SEM micrograph of composite powder containing 1 wt.% of Al_2O_3

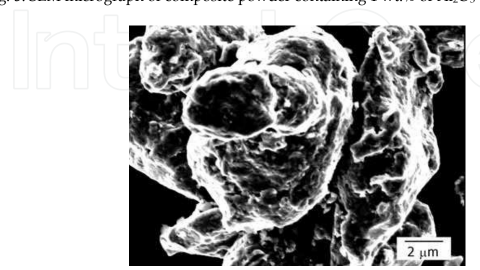


Fig. 6. SEM micrograph of composite powder containing 1.5 wt.% of Al_2O_3

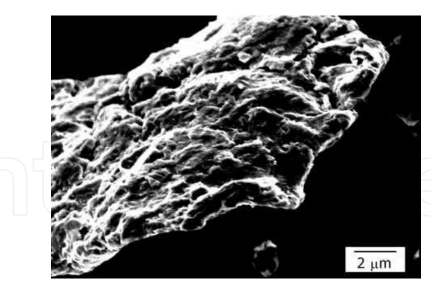


Fig. 7. SEM micrograph of composite powder containing 2 wt.% of Al₂O₃

From these Figures it is obvious that with increased Al_2O_3 content increases particle size. Average particle size of composite powders containing 1% Al_2O_3 was $6\mu m$, 1.5% $8\mu m$ and with $2\%Al_2O_3$ around $12\mu m$. Also, these Figures show that throughout the process of mechanical alloying starting copper particles got reduced in size. Deformation strengthening induced by consecutive plastic deformations by milling media, makes previously ductile particles brittle, further leading to their breakage due to fatigue.

X-ray diffraction patterns (Figure 8.) of powders containing 1, 1.5 and 2 wt. % Al₂O₃ show only peaks corresponding to copper, due to the small amount of Al₂O₃ in system.

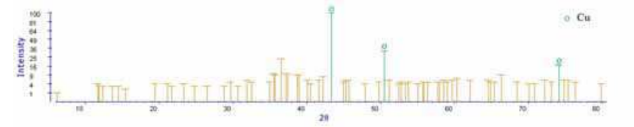


Fig. 8. XRD pattern of Cu-2Al₂O₃

3.2 Compacts characterization

SEM analysis of compacted powders after mechanical alloying containing 1, 1.5 and 2 % of Al_2O_3 sintered at 875°C for 1h is presented in Figures 9-11.

From the presented SEM analysis it is confirmed that with increase of alumina content grain size increases from 1µm for Cu-1Al₂O₃ up to 15-20µm for Cu-2Al₂O₃. Also, sintered compacts exhibit annealing twins and sub-grain boundaries for all three compositions. Twinning might have occurred during the high temperature sintering stage.

Conditions for twins formation are achieved when large number of obstacles is formed in crystal, blocking dislocation movement. Dislocations are accumulated on obstacles causing increase of internal strain in local areas, which together with external strain induce twins formation. Presence of twins indicated lower mobility of dislocations, in other words stabilization of dislocation structure, which is primary condition for improvement of mechanical properties of dispersion strengthened materials.

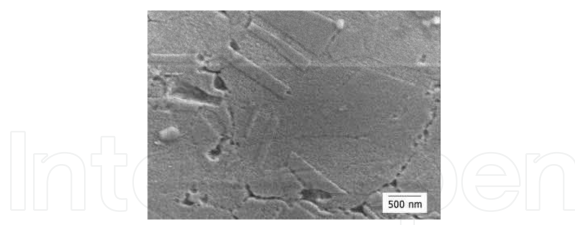


Fig. 9. SEM micrograph of Cu-1Al₂O₃

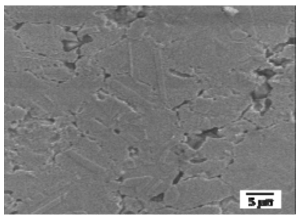


Fig. 10. SEM micrograph of Cu-1.5Al₂O₃

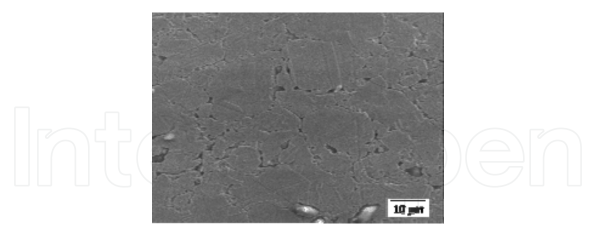


Fig. 11. SEM micrograph of Cu-2Al₂O₃

Size of sub-grains similar to ones presented in Figures 9-11 are expected to be preserved under the influence of temperature as a result of dispersed Al₂O₃ particles blocking sub-grain boundaries and increasing recrystallization temperature, rendering these materials suitable for exploitation at elevated temperatures.

Also, during the period of sintering, formation of a third phase is expected (Lianga et al., 2004), due to the thermodynamically possible eutectic reaction of ($Cu+Cu_2O$) with Al_2O_3 at the contact surface. Through this reaction eutectic tends to expand and reacts with Al_2O_3

forming $Cu_xAl_yO_z$ phase, compatible with both phases at the interface (Yoshino & Shibata, 1992; Jena et al., 2001). The formed third phase influences the dislocation structure resulting in improvement of mechanical properties, whereas good electrical conductivity is retained. According to literature (Yi et al., 1999) chemical formula of this compound might be derived as $CuAlO_2$ or $CuAlO_4$.

Presence of this phase in the structure and especially at the interface could impede crack propagation and result in higher interface fracture energy (Reimanis et al., 1997). Figure 12 presents EDS analysis of produced composites with 1, 1.5 and 2 wt. % Al₂O₃.

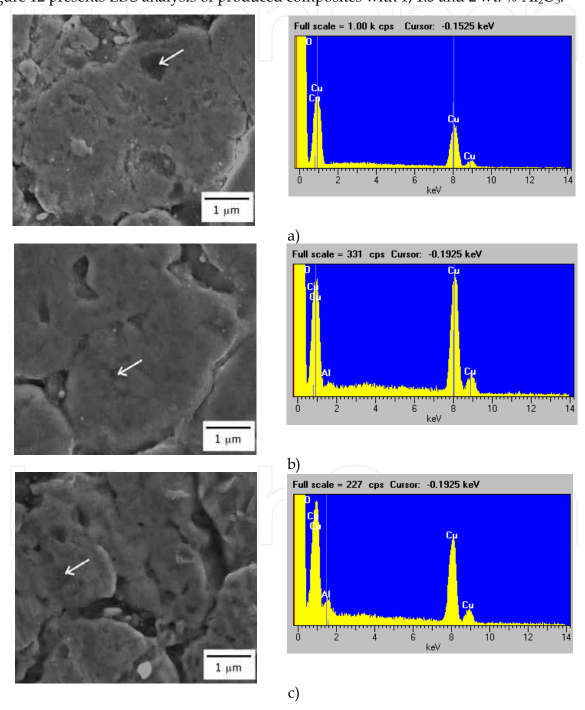


Fig. 12. EDS analysis of marked spots in SEM micrographs of Cu-Al $_2$ O $_3$ composite with a) 1; b) 1.5 and c) 2wt.% Al $_2$ O $_3$

Presented EDS results show that in the composites with $Cu-1Al_2O_3$ aluminum was not detected in the structure, indicating that the whole quantity of Al_2O_3 is incorporated in the copper matrix. With increase of Al_2O_3 content, aluminum is detected which suggests occurrence of segregations in the structure and formation of aluminum-rich regions. These results suggest that the new synthesis method is more suitable for composite materials containing maximum 1wt. % Al_2O_3 . This amount could be fully incorporated in the base-metal matrix playing a role as an obstacle to dislocations, grain and sub-grain boundaries movement. However, this role is diminished with increase of Al_2O_3 content by formation of aluminum-rich regions. Furthermore, Figure 12 shows better compaction for sintered compacts with lower Al_2O_3 content, which is supported by porosity measurements presented in Table 1.

Sample	Min, µm	$M\alpha x, \mu m$	Mean, μm	Std Error	Std Dev.	Vv, %
Cu-1Al ₂ O ₃	0,23	2,41	1,21	0,06	0,43	7,638
Cu-1.5Al ₂ O ₃	0,18	4,46	1,34	0,10	0,72	7,835
Cu-2Al ₂ O ₃	0,45	4,32	1,41	0,11	0,70	8,561

Table 1. Statistical data for the porosity measurements for the samples with different Al_2O_3 content sintered at 875 ° C/1h

These results are also confirmed by measurements of hardness and electrical conductivity as presented in following tables and figures.

Results of HRF hardness measurements are presented in Table 2.

T, °C	725	775	825	875	925		
Cu-1Al ₂ O ₃							
15	52	49	50	52	42		
30	52	54	50	50	52		
60	53	46	48	50	46		
90	48	52	44	45	40		
120	47	46	46	52	44		
Cu-1.5Al ₂ O ₃							
15	42	42	40	42	40		
_30	48	44	40	46	38		
60	44	44	42	40	42		
90	42	43	39	42	38		
120	44	39	40	39	43		
Cu-2Al ₂ O ₃							
15	42	29	22	26	19		
30	26	27	12	20	19		
60	30	24	22	26	20		
90	24	26	15	20	16		
120	30	26	13	23	22		

Table 2. HRF hardness of sintered Cu-Al₂O₃ composites

Results show that with increase of temperature and time of sintering, HRF values slightly and unevenly decrease. Increase of Al_2O_3 content has significant effect on hardness. It was expected that increase of Al_2O_3 content in structure would result with an increase of

hardness, but the results show an opposite trend. With increase of Al_2O_3 from 1 to 1.5 wt.% there is a slight decrease in HRF values, while increase up to 2 wt.% Al_2O_3 provokes a rather significant decrease in HRF. Possible reasons for this behavior is ascribed to a positive effect on dispersion strengthening of the copper matrix which occurred when amount of Al_2O_3 dispersoide is up to 1 wt.%, whereas further increase of Al_2O_3 content has a negative effect on hardness as previously postulated by EDS analysis (Fig. 12). This problem supposedly may be solved using hot extrusion process.

Electrical conductivity measurements of composites with different Al₂O₃ content at various sintering conditions are presented in Table 3.

T, °C t, min	725	775	825	875	925	
Cu-1Al ₂ O ₃						
15	54,88	59,90	56,83	55,05	55,19	
30	53,94	55,39	56,48	57,16	56,06	
60	55,44	54,61	56,28	56,69	55,51	
90	58,00	55,13	54,54	55,25	54,81	
120	56,37	54,24	55,15	56,43	55,87	
Cu-1.5Al ₂ O ₃						
15	42,76	42,49	43,51	43,86	43,81	
30	45,07	42,52	42,51	44,02	47,81	
60	40,94	43,27	45,48	49,59	49,88	
90	39,86	41,87	42,63	46,72	48,08	
120	43,19	39,11	42,76	45,80	48,80	
$\text{Cu-2Al}_2\text{O}_3$						
15	32,66	32,54	30,71	30,35	31,25	
30	29,25	31,27	29,92	31,13	40,70	
60	31,93	30,89	31,72	36,94	39,99	
90	31,35	31,72	31,76	37,61	39,35	
120	30,53	31,66	31,66	33,43	41,21	

Table 3. Electrical conductivity of sintered Cu-Al₂O₃ composites, %IACS

With increase of Al₂O₃ content in composite materials values of electrical conductivity decreases.

Values of electrical conductivity are over 55% IACS for compacts with 1 wt% Al₂O₃, which correspond to a dispersion strengthened copper alloys intended for application at elevated temperatures. The limitation value for application is 50% IACS (Grant et al., 1984).

Measured values of harness, as well as electrical conductivity are highly dependant on Al_2O_3 content and not so much on sintering conditions.

3.3 Characterization of samples after thermo-mechanical treatment

After consolidation of the obtained powders, the compacted samples were uniaxially compressed by cold rolling, reduction degree up to 30%. In order to determine the stability at higher temperatures, the rolled samples were annealed at temperature of 800°C in one hour in the hydrogen atmosphere.

Cold rolling of samples was performed in two steps: first 15% of size reduction and second of 30%. During cold rolling samples with 1.5 and 2 % of Al₂O₃ were degraded, due to inhomogeneous structure and distribution of Al₂O₃, as presented in Figure 12 and increased

porosity with increase of Al_2O_3 (Table 1). Insufficient packing caused formation of cracks in the middle of the sample and disabled sample structure to endure forces of cold plastic deformation.

Considering the presented research results, shown also in (Korać et al., 2010), which suggest that the best combination of mechanical and electrical properties was achieved in the systems with 1wt.% Al_2O_3 sintered at $875^{\circ}\text{C}/1\text{h}$, so further experimental results will be related to these compacts only.

SEM analysis of samples after two stages of size reduction and after heat treatment (HT) is presented in Figure 13.

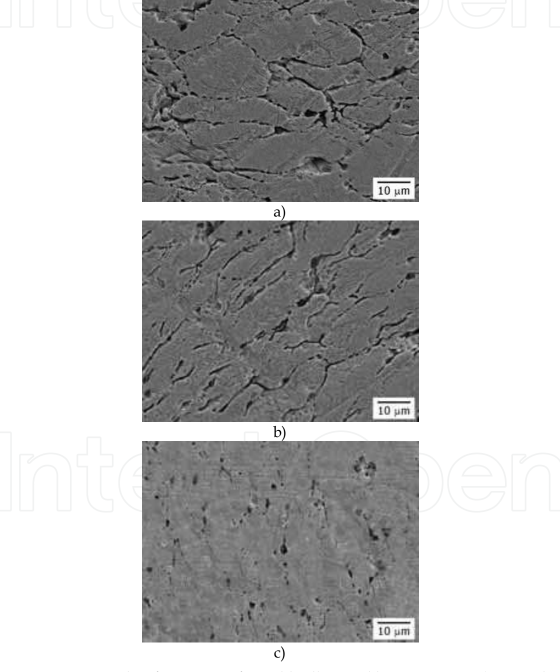


Fig. 13. SEM micrographs of compacts after cold rolling and heat treatment a) size reduction 15%, b) size reduction 30% and c) heat treated

Statistical data on porosity of samples after cold rolling and heat treatment are presented in Table 4. Data for sintered samples is also incorporated in this Table for comparison.

Sample	Min, µm	Mαx, μm	Mean, μm	Std Error	Std Dev.	Vv, %
sintered	0,23	2,41	1,21	0,06	0,43	7,638
15%	0,36	2,84	0,91	0,07	0,490	7,01
30%	0,29	2,14	0,69	0,04	0,337	5,52
HT	0,16	1,56	0,55	0,03	0,29	4,507

Table 4. Statistical data for the porosity measurements for the samples with different Al_2O_3 content sintered at 875 ° C/1h after cold rolling and heat treatment

In the structure of the samples subjected to cold plastic deformation performed by rolling, porosity is also relatively evenly distributed, with a clearly visible porosity of the directed orientation.

The comparative analysis of the porosity testing results of the samples subjected to cold plastic deformation by rolling and the sintered samples indicates the significant porosity in the sintered samples both from the aspect of the pore size, and also from the aspect of their volume share. In the cold deformed samples by rolling, it is clearly noticeable the accumulation of the pores at grain boundaries, which indicates that the pores in the moment of contact with the grain edge during its movement have been too large to be absorbed by the grain edge and disappeared in it, which has resulted in stopping the grain edge to move and its growth.

Rolling can change the distribution and the size of secondary phase, as well as to reduce the number of pores. After rolling, the particles of Al₂O₃ become more dispersive, while the relative density increases (Korać et al., 2008).

The porosity in the structure of the samples subjected to heat treatment after cold plastic deformation is relatively evenly distributed. The pores are arbitrarily oriented, and the shape of the pores is irregular. Based on relative distribution of porosity and statistical data, the size of pores is in the range from 160nm to 1.56 μ m with medium diameter of 550nm. The volume share of porosity is 4.5%.

The comparative analysis of the results of the porosity tests on the sintered samples, the samples subjected to cold plastic deformation by rolling and the HT samples points to significant porosity of the sintered samples and the samples subjected to cold plastic deformation by rolling in relation to the annealed samples as from the aspect of the pore size, so from the aspect of their volume share.

Results of hardness measurements in function size reduction and heat treatment for selected sintering temperature of 875°C are presented in Fig. 14. and for electrical conductivity in Fig.15.

From Fig.14 it could be noticed that with increase of size reduction hardness values are increasing up to the level of 90 HRF indicating stabilization of dislocation structure. Also, curves of dependence HRF-time are stabilizing. Stabilization of curve could be the result of structure compaction and reduction of pore volume fraction in structure. After heat treatment harness values decrease up to the value of 55 HRF. Change of curve slope could be detected for sintering time of 60min. After 30% reduction at 60min slope is changing into plateau. Heat treated samples exhibit slight decrease of hardness values after 60min.

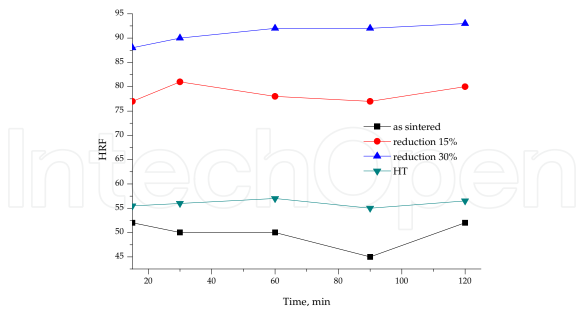


Fig. 14. Dependence of HRF hardness from size reduction and heat treatment for selected sintering temperature of 875°C

Values of electrical conductivity for as sintered samples are over 55% IACS, which correspond to dispersion strengthened copper alloys intended for application at elevated temperatures. With increase of size reduction electrical conductivity is increasing up to the 59.5 %IACS. After heat treatment electrical conductivity continues to increase up to 60.75%IACS for sintering time of 60min, when plateau is reached. After 60 min of sintering hardness and electrical conductivity both reached constant values, indicating completed stabilization of structure.

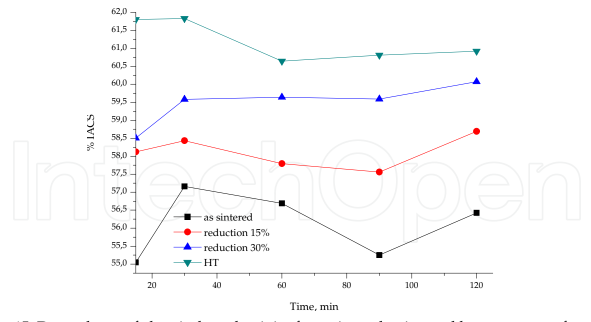


Fig. 15. Dependence of electrical conductivity from size reduction and heat treatment for selected sintering temperature of 875°C

Aiming to evaluate the stability of structure for the sintered samples and each phase of the thermo-mechanical treatment process, the examinations on tribological properties were performed.

It is well known that the HT metals with the surface centered cubic crystal structure have significantly higher resistance to erosive wear in comparison to metals with the spatially centered cubic crystal structure of similar hardness. Cold rolling prior to wear testing does not bring significant improvement in resistance to erosive wear. Also, reducing the speed rate of the erosive wear is achieved with an increase in the index of the deformation hardening. Increasing in the index of the deformation strengthening effects on increasing the value of critical voltage that is required to starting with localize the deformation in the process of wear (Liu et al. 2007).

Tribological tests have shown that the best resistance to wear have the samples after heat treatment (Fig. 16), which can be explained by the fact that after annealing recrystalization and characteristic growth of the grain do not occur, due to block in movement of the grain boundaries homogeneously distribution of alumina and formation of the third phase, identified in (Jena et al., 2004; Jena et al., 2001; Lee & Kim, 2004, Anđić, 2007, Korać, 2009). This is confirmed by the microstructural examinations of the analyzed samples.

Based on the results of quantitative analysis and tribological examinations, it can be concluded that the effect of porosity has a significant influence on the tribological properties, that is, as the volume share of porosity reduces as tribological properties improve.

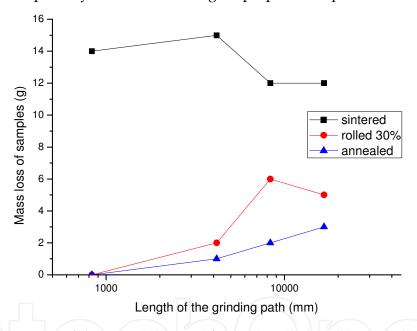


Fig. 16. Wear resistance of the sintered (875°C/60min), rolled and annealed sample

The significant influence on the tribological properties has multiple strengthening mechanisms of the analyzed systems schematically presented in Figure 17.

In the observed system, strengthening the base of copper is achieved by dispersion strengthening, due to the dispersion of fine particles of Al_2O_3 in the base, strengthening by grain boundaries, because of the third phase appearance, as well as the deformation strengthening and strengthening by annealing. During mechanical alloying the coating copper particle with the particles of the composite 50% Al_2O_3 occurs, which is built into its surface. Along with the process of sintering occurs the formation of the compact structure and formation of the third phase on the grain boundary, which causes strengthening on the grain boundaries. Due to the plastic deformation occurs the deformation strengthening, and after heat treatment occurs the strengthening by annealing, which is confirmed by the results of the tribological examinations.

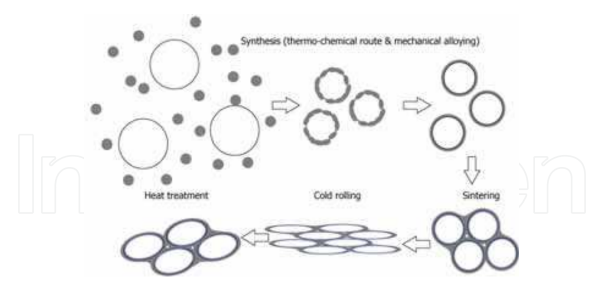


Fig. 17. Proposed multiple strengthening mechanisms

4. Conclusion

Synthesis of nanocomposite Cu-Al₂O₃ powder suitable for obtaining multiple strengthened systems could be successfully performed by combination of the thermo-chemical procedure and the mechanical alloying. To optimize the whole scope of properties that are expected from these materials, the combination of thermo-chemical procedures and the procedure of mechanical alloying is a completely new approach to the synthesis of powders. Thus obtained powders allow obtaining the final product with the excellent effects of strengthening.

Presented results show possibility of usage composite powders based on copper and alumina synthesized by novel method for production of compacts with suitable properties for exploitation at elevated temperatures, i.e. increased hardness with electrical conductivity appropriate for oxide dispersion strengthened copper alloys intended for application at elevated temperatures. Achieved values after thermo-mechanical treatment for hardness are 57HRF and for electrical conductivity 61%IACS.

Limitation for dispersoide content is 1wt. %, due to the decomposition of compacts with higher Al₂O₃ content during mechanical processing. Increased Al₂O₃ content could be achieved by using hot extrusion process.

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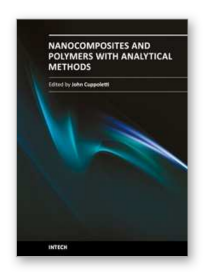
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