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Structural, Mechanical, and Tribological Characterization of Magnetic Pulse Compacted Fe–Cu Bimetallic Particles Produced by Electric Explosion of Dissimilar Metal Wires

Alexander Pervikov 1, Anton Khrustalyov 1,2, Andrey Filippov 1, Yuri Mironov 1, Aleksander Lozhkomovev 1, Marat Lerner 1 and Sergei Tarasov 1,*

1 Institute of Strength Physics and Materials Science SB RAS, 634055 Tomsk, Russia; pervikov@list.ru (A.P.); tofik0014@mail.ru (A.K.); avf@ispms.ru (A.F.); myp@ispms.ru (Y.M.); asl@ispms.ru (A.L.); lerner@ispms.ru (M.L.)
2 National Research Tomsk State University, 634050 Tomsk, Russia
* Correspondence: tsy@ispms.ru

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Abstract: Bimetallic 73 wt.% Fe–Cu nanoparticles have been produced using electric explosion of two immiscible metal wires and then consolidated into disks using magnetic pulse compaction. The compacted disks have been characterized for phase composition, mechanical strength, and high-temperature steel ball-on-disk sliding friction. The sample possessed good flexural and compression strength. Friction and wear reduction were observed during sliding test at 400 °C, which was explained by intense tribosynthesis of cuprospinel CuFe2O4 nanoparticles, which served to reduce adhesion between the ball and disk.

Keywords: electric explosion; bimetallic particle; pseudoalloy; cuprospinel; triboxidation

1. Introduction

The interest in preparing and using Fe–Cu pseudo-alloys started since the 1940s [1]. One of the potential applications was improving the wear resistance of materials used in tribotechnical devices. The sintered iron–copper alloys were used as electrotechnical material for asynchronous motors since they possessed optimum magnetic permeability and specific electrical resistance [2]. Consolidated Fe–Cu composites [3] containing from 2 to 40 mass% Cu had higher mechanical strength as compared to pure iron ones, despite the fact that their tribological behavior was almost similar to that of pure iron composites. The Fe–Cu composites containing more than 10 to 70 mass.% Cu demonstrated improved corrosion resistance and tribological characteristics comparable to those of anti-friction bronzes.

One of the feasible applications for Fe–Cu composites is automotive brake pads and the tests show a stable friction coefficient as well as high durability during tribological tests [4]. Also, Fe–Cu composites served as the matrix for making multicomponent brake pad materials by filling it with various non-metal additives such as graphite SiO2, ZrO2, etc. [5]. The effect of copper content on tribological characteristics of Fe–C–Cu composites was studied by Dyachkova et al. [6]. It was shown that friction was reduced for the low-copper composites under constant load and increased for the incrementally increased load.

The effect of adding copper nanosized powders to lubricant oil on friction is known to reduce friction and form an iron-oxide layer on the steel surface [7]. According to XPS data, copper particles were found in this layer in a metallic form so that the mechanically mixed layer consisted of FeO3...
and metallic copper. More intense mechanical alloying of Cu–Fe powders resulted in formation of FCC Cu(Fe) and BCC Fe(Cu) solid solutions as well as CuO and Cu₂O [8].

However, tribological behavior of the Fe–Cu matrix has not been fully elucidated. Structural adaptability of materials to friction is determined by their structural as well as mechano-chemical evolution under severe sliding friction conditions. It is especially true for high-temperature sliding friction when tribooxidation becomes very intense and both components may oxidize instead of only iron as it happens at a low sliding speed. Tribooxidation and tribosynthesis of mixed oxides may occur, which is known to be an adaptive mechanism to reduce both wear and friction under high temperature sliding friction [9]. Practically immiscible metals such as Cu and Fe can form neither alloy nor chemical compound but instead may form mixed oxides known as delafossite CuFeO₂ and cuprospinel CuFe₂O₄.

Bimetallic nanoparticles draw great attention of scientists during past decades “as an emerging new class of materials” that allows not only to combine the characteristic of both dissimilar metals, but also obtain some new unique characteristics [10]. A great variety of their shapes and structures may be obtained when using different production processes such as electric explosion of wires, mechanical alloying, chemical, and electrochemical methods. Electric explosion of intertwined wires (EEIW) allows producing the oxide-free bimetallic particles and, therefore, it is the preferential method for obtaining electro conductive materials [11,12]. It is also a necessary condition for synthesizing the dissimilar immiscible metal bimetallic particles that this method provides a very high cooling rate at the level of ~10⁷–10⁹ K/s [13,14]. An electric explosion of wire (EEW) phenomenon is observed when a 5–15 mm length and 0.1–0.5 mm diameter conducting wire is energized by an electric current with pulse duration 0.5–5.0 μs and density as high as /cm² is thus causing its atomization. Such a process is used to obtain metallic nanoparticles of metals and alloys in either Ar or He atmosphere at 10⁵–5 × 10⁶ Pa. The explosion product expansion velocity of the buffer gas is at the level of 10³ m/s [11].

Improving the mechanical strength of the composites is connected with refining the composite grains by taking advantage of compacting the bimetallic nanoparticles and using a compaction process that allows preserving their size.

When using the bimetallic nanoparticles, which are composed of two dissimilar metals in a singular particle, for obtaining a composite, it is reasonable to employ some dynamic compaction methods in order to avoid grain growth and delamination. Therefore, more homogeneous distribution of components will be provided during compaction. One of the potential methods is the magnetic pulse compaction (MPC), which is based on transformation of an electric energy pulse into a mechanical powder compacting impact [15–19]. The impact rise time is about hundreds of microseconds, which provides compaction pressure in the range 1–5 GPa and the powder particle displacement rate in the order of 10–100 m/s. It should be noted that that high velocities, which in fact are several orders of magnitude higher than ~10⁻³ to 10⁻⁴ m/s achieved in static pressure compaction, are necessary to prevent the particle coalescence growth. At the same time, the particle velocities achieved during MPC are at least one order of magnitude lower than that of the soundwave in metals so that no shock wave is generated in the compacting powder, which could result in local cracking in the tensile stress regions.

The objective of this work is to study the EEW Fe–Cu particles, use them for magnetic pulse compacting, and characterize the compacted composites for mechanical strength and high-temperature tribological behavior.

2. Materials and Methods

The bimetallic Fe–Cu powder particles were produced using electric explosion of a twisted pair of dissimilar metal wires in a chamber filled with argon at 3 × 10⁵ Pa. More detailed description of the EEW process is provided elsewhere [20]. The EEW machine functioning is based on using an RLC-circuit with parameters, as follows: $L \approx 0.75 \mu$H, $C = 3.2 \mu$F, $U_0 = 29$ kV, where $L$ is the inductance, $C$ is the electric capacity, and $U_0$ is the charging voltage (Figure 1). A capacitor battery (C) is charged from the direct current power source (PS) until reaching $U_0$ voltage level. On reaching the $U_0$, the air-
gap discharger is activated, and the circuit is energized by a current pulse of $10^7$ A/cm$^2$ density. The wires suffer burst atomization into fine drops.

![Circuit diagram of the machine employed for the production of Fe–Cu nanoparticles.](image)

Both wires had the same length of 75 mm and different diameters of $\varnothing$0.34 mm (Fe) and $\varnothing$0.2 mm (Cu). Their sublimation energies were 418 and 112 J, respectively [21]. The wire diameter ratio resulted in EEIW producing the powder composed of 73% wt. Fe and 27% wt. Cu.

The time dependencies of the EEIW current $I(t)$ and voltage $U(t)$ inherent for iron and copper were determined using a TDS2022B oscilloscope (Tektronix Inc., Beaverton, OR, USA) using a scheme shown in our previous paper [22]. The input energy $E(t)$ levels were calculated using Equation (1), as follows:

$$E(t) = \int I(t)U_r(t)dt,$$  \hspace{1cm} (1)

The $U_r(t)$ dependence was then obtained using Equation (2);

$$U_r(t) = U(t) - U_l(t),$$ \hspace{1cm} (2)

where $U_r(t)$ and $U_l(t)$ are the resistive and inductive components of the voltage drop on an electrical circuit section, which includes the wire, respectively. The inductive one was calculated using Equation (3):

$$U_l(t) = L \frac{dI}{dt},$$ \hspace{1cm} (3)

where $L$ is the inductance of the wire-containing circuit, 0.22 μH, and $dI/dt$ is the EEW current derivative.

The resulting powder was characterized for particle size distribution and particle microstructure using a TEM instrument JEOL JEM-2100 (JEOL Ltd., Akishima, Japan) and X-ray diffractometer XRD-6000(Shimadzu Corp., Kyoto, Japan), CuK$\alpha$ wavelength. Samples for TEM were prepared by dispersing 50 mg powder in 100 mL alcohol and sonication for 5 min in order to provide the powder deagglomeration. Then, a 10 μL portion of the resulting suspension was placed on a carbon-coated $\varnothing$3 mm gold mesh substrate and dried.

The powder phase composition was determined using PDF 4+ databases and full-profile analysis software POWDER CELL 2.4 (Federal Institute for Materials Research and Testing, Berlin, Germany).

The particle size distribution was reproduced using the data obtained from sedimentation in a disk centrifuge DC24000 (CPS Instrument Inc., Prairieville, LA, USA). The resulting sonicated for 5 min suspension contained 20 mg particles dispersed in 10 mL alcohol. The duration of analysis was
30 min. The particle size distributions were obtained by counting no less than $4.386 \times 10^9$ isolated particles.

The mean particle size ($a_s$) was determined from measuring the BET specific surface area in a Sorbtometr-M (Catakon Ltd., Novosibirsk, Russia) instrument and formula

$$a_s = \frac{6}{\rho S}$$

where $\rho$ is the density (~8 g/cm$^3$); $S$ is the specific surface area, m$^2$/g.

To produce solid samples, the EEIW bimetallic powders were subjected to magnetic pulse compaction (MPC) at the Institute of Electrophysics UrO RAN, (Yekaterinburg) [23]. The total amount of powder intended for the compaction was 400 g. A portion of 32 g of powder was loaded into a mold [23] and then pre-compacted in a static mode until reaching about 40% of the theoretical density. Then, it was degassed at 350 °C and 1 Pa for 4 h prior to magnetic compaction, which was carried out at electric capacity and voltage values 2.2 mF and 4.1 kV, respectively. The MPC parameters such as above have been selected experimentally by achieving the consolidated sample compression strength and density. The compacted sample density was determined using the hydrostatic weighing. On magnetic compaction, the resulting samples were annealed in vacuum at 450 °C for 2 h to relieve the macrostresses induced during compaction. The compacted samples had 5.94 mm height and 30.5 mm diameter.

Microstructural evolution of the compacted samples was examined using an SEM instrument Quanta 200 3D attached with an equipment for ion milling.

Mechanical tests for compression and flexural strength were carried out using an Instron 1195 (Instron, USA) tensile machine at loading rates 50 $\mu$m/min and 100 $\mu$m/min for flexural and compression tests, respectively. The compression and flexural test samples had their dimensions as follows: 10 mm diameter, 5 mm height and $5 \times 5 \times 30$ mm$^3$, respectively.

Microhardness tester Buehler 1600-6400 (Buehler, Germany) was used to determine the microhardness numbers at 25 g and 50 g loads. At least three samples of each type were tested.

Sliding friction experimenting was carried out using a high-temperature nanotribometer THT-S-BE-0000 (CSM Instruments, Peseux, Switzerland) under test conditions as those shown in Table 1.

| № | Temperature, °C | Rotation rate, RPM | Radius, mm | Time, s | Linear speed, m/s | Normal load, N |
|---|-----------------|--------------------|------------|--------|------------------|----------------|
| 1 | 25 | 300 | 8 | 3600 | 0.25 | 10 |
| 2 | 100 | 300 | 8 | 3600 | 0.25 | 10 |
| 3 | 250 | 300 | 8 | 3600 | 0.25 | 6 |
| 4 | 400 | 300 | 8 | 3600 | 0.25 | 5 |

3. Results

3.1. The EEIW Fe–Cu Nanoparticle Synthesis and Characterization

The dissimilar metal EEIW was a fast-occurring process, which was characterized by fast changes in the process parameters such as current, voltage, and input energy belonging to different wires (Figure 1). It was demonstrated (Figure 2) that copper wire was the first to explode at the moment of time $t$~$1.3 \mu$s while the iron one exploded a bit later at $t$~$1.8 \mu$s. The input energy values corresponding to the moment of explosion were 170 J and 450 J for copper and iron wires,
respectively. These energy values related to corresponding sublimation energy values for copper and iron as $1.5 \times E_s$ and $1.1 \times E_s$. The next stage of the EEIW was characterized by forming an arc discharge in the explosion products at $t \sim 2.45 \mu s$. It is seen from $I_{Cu}(t)$ and $I(t)$ curves (Figure 2) that the arc stage current flowed mainly via the copper wire explosion products, thus extra heating and even evaporating them [24].

Both theoretical and experimental studies show that nanoparticles are formed during EEW by means of condensed cluster coagulation in the expanding EEW products [25–28]. These clusters are generated during EEW as a result of inhomogeneous energy release at the crystallite boundaries and instabilities caused by such an overheating so that the metal becomes fragmented into the nanosized clusters [29–31].

![Figure 2. The EEIW current $I(t)$, voltage $U(t)$ and input energy $E(t)$ during double-wire explosion.](image)

The resulting EEIW Fe–Cu nanoparticles were spherical (Figure 3a) and demonstrated the particle size distribution (Figure 3b). The EDS mapping shows that each particle is composed of copper and iron (Figure 3c,d). That type of particle could be identified as so-called Janus particles when one metal part maybe eccentrically encapsulated inside the other [32] or two dissimilar metal parts welded. The mean by five-measurement BET specific surface area of the powder was $9.1 \pm 0.9 \text{ m}^2/\text{g}$ so that according to formula 4, the mean particle size was $a = 82 \pm 9 \text{ nm}$. This value is close to that obtained from the TEM analysis.

The SEM image in Figure 4 shows that EEIW powders contained some amount of micron-sized large particles along with the nanosized ones, which plausibly formed when the energy input reached the level of the wire metal sublimation energy. It is known [33,34] that the micron-sized particles may form in such a case amidst the EEIW expanding products. However, the percentage of that large particles is about two orders of magnitude lower than that of nanosized ones.
Figure 3. TEM images of the bimetallic EEIW Fe–Cu particles (a,c), particle size distribution (b), (d) the EDS Fe–Cu map of the particles shown in Figure 3c.

Figure 4. The SEM image of the as-synthesized Fe–Cu micron-sized particles.

Phase composition of the Fe–Cu powder was analyzed using the XRD. The diffractogram in Figure 5 shows the presence of α-Fe, γ-Fe, Cu, and CuO. Lattice parameters of phases were close to the standard values.
3.2. Compacted Sample Characterization

The as-compacted samples had 93.8% density of the theoretical one as measured using the hydrostatic weighing. The optical microscopy image in Figure 6 and SEM images and EDS profiles in Figure 7a,b show that, microstructurally, sample was composed of very fine copper and iron crystallite mixture as well as isolated coarse iron and copper particles. These coarse particles were less oxidized as compared to the fine mixture (Figure 7a,b). As follows from the XRD pattern in Figure 8, the lattice parameters of both metals differed from their standard values. Also, the presence of CuFe2O4 (Fd3m, a = 8.394 Å) and FeO (Fm3m, a = 4.29 Å) oxides was detected.

Figure 5. The XRD pattern of the as-synthesized Fe–Cu powder.

Figure 6. Microstructure of a magnetic pulse compacted sample.
Figure 7. The SEM images of as-compacted samples with the imposed EDS profiles of oxygen (red line), iron (green line), and copper (blue line) showing large iron particles (a) and fine copper/iron crystallite mixture (b).

Figure 8. The XRD pattern of as-compacted samples.

Taking into account dynamic mode of compaction, it was suggested that the compacted samples might contain some residual stresses and therefore special studies using the known \( \sin^2 \psi \) method were undertaken. The results are shown in Figure 9. When comparing the lattice parameters from the approximation lines for \( \sin^2 \psi = 0 \) and \( \sin^2 \psi = 1 \), i.e., measured along the normal to the surface, to those obtained from the symmetrical scheme, one can see that strain values were very low, so that almost no residual stress was induced during the magnetic pulse compaction.
Figure 9. Approximation lines for lattice parameters of Cu (red line) and Fe (black line) as measured from an asymmetrical scheme according to the \( \sin^2\psi \) method. The dotted lines show corresponding lattice parameter values obtained from a symmetrical scheme.

3.3. Mechanical Strength

Mechanical strength of the as-compacted samples was determined using flexural and compression tests. The corresponding flexural strength was 880 MPa. Compression yield stress and ultimate stress were 765 MPa and 660 MPa, respectively (Figure 10a,b). The microhardness was Hv = 390 \( \pm \) 20 kg/mm\(^2\).

Figure 10. Flexural (a) and compression (b) stress–strain diagrams for as-compacted samples.

The SEM in Figure 11 demonstrates the morphology of the fracture surfaces after flexural test. It is seen from this image that fracture surface was represented mainly by ridges formed during rupturing of plastically deformed and flattened grains.
3.4. Tribological Test

Wear tracks of the samples tested at different temperatures were examined using optical confocal and scanning electron microscopies. Wear tracks of samples tested at 25 °C appeared dark and 100 °C appeared as a light-grey color. Wear tracks of the sample tested at 250 °C had a black color, and finally, the sample tested at 400 °C acquired a brown color.

Temperature dependencies of friction coefficient \( f \), sample wear \( W_{\text{disk}} \), and steel ball wear \( W_{\text{ball}} \) are shown in Figure 12. All these parameters demonstrate qualitatively similar behavior, achieving their maximum values at 250 °C and then reducing at 400 °C.

Wear tracks formed on the samples in tribological testing at different temperatures were examined and measured to obtain their cross section areas (Figure 13), and the calculated total wear \( W_{\text{disk}} \) (Figure 12) related to friction force work in order to take into account the fact that normal load was about two times lower for 250 °C and 400 °C tests. Wear of the steel ball counterbodies \( W_{\text{ball}} \) was determined as the wear spot surface area projected on a plane since real wear spot surface was not a flat one (Figure 14).
Figure 12. Tribological parameters such as friction coefficient ($f$), disk sample wear ($W_{\text{disk}}$), and the ball wear ($W_{\text{ball}}$) vs. the test temperature.

Figure 13. Cross section areas of wear grooves formed on the sample disks in sliding friction by steel balls at different temperatures.

Figure 14. The wear spot surface profiles on the steel balls.
The SEM images of wear track surfaces are shown in Figure 15. All the worn surfaces were composed of flat and smooth bearing areas and rough non-contacted areas with wear debris. In addition to structural changes, the wear tracks of the wear track material experienced mechanochemical changes. The EDS profiling the worn surfaces allowed us to show that tribooxidation degree increased with the temperature. Such a finding is illustrated by the data presented in Figure 16 by the examples of Fe/O and Cu/O EDS concentration ratios. The higher test temperature led to the diminishing of both ratios. This means that the degree of oxidation increased with the test temperature.

Figure 15. The SEM images of wear track surfaces on samples tested at: (a) 25 °C, (b) 100 °C, (c) 250 °C, (d) 400 °C.
Figure 16. Copper/oxygen and iron/oxygen atomic concentration ratios on the wear track metal depending on the test temperature.

It is a well-known fact that the accuracy of EDS with respect to detecting light elements is rather low, but that would be enough for comparing two oxides.

The XRD phase composition of the worn tracks is shown in Figure 17. One can see that CuFe₂O₄ peaks became more intense with the test temperature, so the corresponding phase contents were calculated and presented in Table 2. The worn surface of the sample tested at 25 °C and 100 °C showed the cuprospinel content at the level of as-compacted sample, i.e., 4.2–4.3 wt.%. However, these values include the same phase contents in the bulk of the samples, which resulted from oxidizing during magnetic pulse compaction. Nevertheless, the most intense synthesis of CuFe₂O₄ occurred during sliding at 400 °C, so even the iron oxide content formed during sliding at 250 °C reduced.

Figure 17. XRD diagrams of the wear track surface of samples tested at different temperatures.
Table 2. Phase composition of the wear track subsurface metal.

| Test temperature | Phase content, wt.% | $\alpha$-Fe | Cu | CuFeO$_2$ | FeO | Fe$_2$O$_3$ |
|------------------|---------------------|-------------|----|-----------|-----|------------|
| Consolidated sample |                    | 58.2        | 37.2 | 2.9      | 1.7 | —          |
| 25               |                    | 61.3        | 34.4 | 4.3      | —   | —          |
| 100              |                    | 60.7        | 35.0 | 4.2      | —   | —          |
| 250              |                    | 59.5        | 31.0 | 6.6      | —   | 2.9        |
| 400              |                    | 58.0        | 31.7 | 8.0      | —   | 2.3        |

In order to clearly see the differences, the XRD patterns were obtained from the as-compacted metal surface on the sample’s bottom and then deduced from those obtained on the worn surfaces. Such an approach allowed for understanding that there is no great difference in phase compositions of samples tested at 25 °C, 100 °C, and 250 °C in terms of CuFeO$_2$ peaks (Figure 18). Only weak and wide remainder CuFeO$_2$ peaks can be seen on the worn surfaces of these samples. Therefore, there was no big difference in CuFeO$_2$ content on these worn surfaces and in the bulk of the samples. On the other hand, the remainder XRD diagram obtained from the worn surface of the sample tested at 400 °C clearly shows the corresponding rather narrow peaks, thus giving evidence in favor of the CuFeO$_2$ tribosynthesis at that high temperature.

Figure 18. The remainder XRD plots showing the difference between the worn surface phase composition and that of the sample back side.

4. Discussion

In distinction to many other methods, the EEW bimetallic Fe–Cu particles were produced using the electric explosion of dissimilar metal wires, i.e., a unique process that allows mixing metals in a liquid state. It is necessary to note that the phase composition of the Fe–Cu particles is sensitive to
the EEW process parameters. In contrast to the presented here results, the bimetallic 72 wt.% Fe-28 wt.% Cu nanoparticles were obtained earlier under conditions of a synchronized HV discharge without an arc stage [35]. It follows from the comparison that increasing the input energy level is accompanied by increasing the content of γ-iron in the resulting powder. According to [36], the presence of metastable phases in the EEW products is the consequence of a high rate cooling. When EEW is carried out with the arc stage, the cooling rate of nanoparticles must be lower than that of particles obtained under EEW without the arc stage and therefore no γ-iron should have been formed. However, the XRD clearly shows its presence (Figure 4). In our opinion, such a finding may be explained by extra stabilization of the FCC lattice from Cu atoms, whose solubility in γ-Fe is about 9 wt.% as compared to 0.4 wt.% in α-Fe. The arc stage provides extra heat that allows better intermixing of Cu and Fe atoms in the nanoparticle and thus more homogeneous distribution of Cu in Fe. On cooling, the FCC γ-iron becomes more stabilized.

The role of copper in increasing the content of austenite was reported by Amran et al. [37]. Another rationale may be that the γ→α transformation is sensitive to the grain size, and nanosized particles acquire extra stabilization at the high-temperature phase [38].

No residual stresses were induced in the compacted metal. However, compaction resulted in oxidizing the metal especially at the grain boundaries. The majority of the oxides is represented by the iron oxide FeO with lower content of high-temperature cubic spinel CuFe2O4. It is known [39] that among many other methods, MeFe2O4 spinels can be obtained via mechanical milling, powder metallurgy, solid state reactions, etc. High local temperatures and pressure are developed during magnetic pulse compaction that serve to the synthesis. It is feasible that a new magnetic pulse process could be developed for fabricating spinels. Cuprospinel CuFe2O4 is a widely applied material that has two structural modifications such as cubic and tetragonal ones. Structural transformation from cubic to tetragonal can be facilitated by applied external pressure [40]. At the same time, the cubic phase is more stable at temperatures above 400 °C.

The MPC samples demonstrated their strength characteristics at the level of those obtained using a cold isostatic pressure compaction of Fe–Cu nanoparticles [35]. In addition, the MPC sample strength characteristics may be improved by reducing the content of oxides. Their microhardness ~390 ÷ 20 kg/mm2 is comparable to that of obtained by high-pressure torsion (~ 420 kg/mm2) [41]. The high mechanical characteristics of the MPC samples indicate high potential of this compaction method for obtaining multi-functional bulk nanostructured materials.

Tribological behavior of the samples shows its temperature dependence. The low-temperature sliding friction test is characterized by low wear. The main wear mechanism is by adhesion and oxidation. Heating the samples to 100 °C resulted in somewhat higher wear of sample and lower wear of the ball. Such a behavior may be explained by thermal softening of the sample while the ball was still hard enough and resisted wear. The maximum wear of both disk and ball was achieved after testing at 250 °C. The wear track on the sample was a deep groove while the ball surface had a hump in the middle. The maximum friction has also corresponded to this temperature.

Further increasing the test temperature led to reducing both friction and wear. Reduction in wear and friction during high-temperature sliding at 400 °C may be provided by mechanochemical formation of extra cuprospinel particles, which were detected using the XRD (Figure 17). Formation of cuprospinel in fretting wear between Incoloy and technically pure copper was observed by Soria et al. [42] during characterization of damage and triboparticles resulting from fretting of Incoloy 800 steam generator tubes against different materials.

It is understandable that the higher test temperature enhances tribooxidation so that both Cu/O and Fe/O atomic concentration ratios reduce and reach their minimum values after testing at 400 °C.

5. Conclusions

The EEW Fe–Cu bimetallic nanoparticles structurally consist of two immiscible metals bonded into a singular particle. That means that the real powder dispersity becomes even higher as compared to that of mono-metallic ones. Such a material requires suitable compaction method, which is implied to preserve a high degree of structural dispersity in a compacted sample as well as provide acceptable
mechanical and functional characteristics. The results of this investigation allow for concluding that magnetic pulse compaction may be a suitable technique for achieving that because it is fast enough to limit the grain growth and preserve the nanostructure. At the same time, it provides acceptable strength of the consolidated samples.

Tribological adaptation mechanisms are based on an idea that two immiscible metals will form a mixed oxide, which will reduce both friction and wear. It was found out that in the case of Fe–Cu nanocomposite, such an adaptation mechanism was represented by formation of cuprospinel during the 400 °C sliding test, which allowed for smoothing the worn surfaces and thus reducing both wear and friction. That low-temperature synthesis is provided by intense and severe subsurface deformation and extra heating the subsurface by friction-generated heat.

Author Contributions: A.P., A.K., Y.M., and A.F. performed sample preparation and characterization, A.L., M.L., and S.T. performed conceptualization, designed the experiments, and analyzed the data. A.P. and S.T. wrote this paper.

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