

The study of the effect of chromium boron (CrB_2) hardening additive on the developed pg-z40 surfacing powder

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Development of a new composite powder alloy to increase wear resistance and impact resistance in gas-powder surfacing of the existing alloy PG-Zh40. Optimization of composition of new composite alloys by an injection 10% of chromium-boron hardening additive and characterization of alloy formation conditions. Phase formation conditions and structure of the new composite surfacing material with boride hardening during mechanical alloying. The following rational part of the new powder filler alloy has been received: Fe = 32-40%; C = 1.1-1.4%; Cr = 14-15%; Si = 2-3%; B = 2-2,9%; Ni = 30-32%, Cu = 2-3% with 0 ÷ 44 HRC hardness.

Keywords: Self-fluxing alloy, gas powder surfacing, durability, mechanochemical technology, attritor, granulation

INTRODUCTION

The world economy annually loses approximately 80 billion USD due to wear and corrosion, but adequate proactive wear protection can help to avoid these losses. This protection involves surfacing of new parts as well as reconditioning and return of worn ones to economic circulation. Surfacing with materials with high performance characteristics is an effective method of machinery parts surfaces hardening. This method is cost-effective, because surfacing is applied only to the surfaces functioning in an aggressive wear environments, and as a rule, the weight of deposited material is very low in comparison with the total weight of a part. Durability of hardened parts is determined by the characteristics of deposited material, and for this reason, the materials or alloys used for surfacing are selected on the basis of the part's operational environment and the surfacing method.

The new self-fluxing surfacing ferrous powder material with hardening additive developed by our team will be used for the reconditioning of components of equipment and machinery operating in abrasion wear, corrosion, high temperature or aggressive environment exposure conditions.

Currently, there are many self-fluxing surfacing powder nickel- and copper-based alloys produced under various methodologies. These alloys have started to take the lead among the materials commercially produced by the world's leading companies such as NACA, JNCO, Battelle, Cabot,

BBC, Vienna, KRUPP etc. Alloys with hardness specified in the range from 35 to 55 HRC (such as PSR-2, PSR-3, PSR-4 etc. (GOST 21.448-75)) were developed to create coating of different hardness. All these alloys are cobalt-, nickel- and copper-based with various carbide-forming additions ensuring required physical and mechanical properties of the surfaced (applied) coating.

The goal of this research is to develop a new alloy technology, based on the previously developed self-fluxing surfacing powder alloy PG-Z40 [1,2], but with the addition of chromium boron hardening additive. The team also studied the formation of structure, phase composition and features of gas-flame coatings. To try out the new alloy technology involving the addition of CrB_2 the team used self-fluxing surfacing powder alloy PG-Z40 with 50-160 μm grading fraction and the following chemical composition: Fe=38%; C=15,1%; Si=3,0%; B=2,9%; Ni=33%.

RESEARCH METHODOLOGY

Fe, Ni, Cr, Si, Cu, B, C powders and CrB_2 hardener were used as initial reagents for the production of the self-fluxing surfacing powder. Copper was added to increase fluidity of the melt and to improve anti-corrosive properties of the alloy. Vanadium was added to increase the alloy's cold resistance; vanadium also forms strengthening phases with carbon, resulting in the improvement of the alloy's wear resistance. The increase of carbon content is necessary for the formation of carbides. Carbides evolve from the liquid solution during gas-powder

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surfacing, become crystallization centers and contribute to structure refinement resulting in the improvement of the alloy's durability.

The composite material was developed under mechanochemical methodology with the use of attritor [3], and the produced powder was granulated with planetary granulator to create powder of required grain size. The structure of the surfacing alloy, its physical and mechanical performance properties were identified with standard techniques. In particular, the structural analysis in the surface layers of the surfacing ferrous alloy was carried out using metallographic analysis as well as scanning and transmission electron microscopy. X-ray crystallography was used to determine the phase composition of samples. Microhardness was measured with PMT-3 microhardness tester using static indentation method in accordance with GOST 9450-76. Surfacing layer adhesive strength was examined with CDM10/91 strength testing machine. Surfacing layer hardness was determined with TP-7R-1 material and alloy hardness measurement device using Vickers method. Technological properties of produced granulated powder mix were examined with the use of gas-flame surfacing on various steel samples.

Phase composition of the new surfacing alloy was determined with D8 Advance (BRUKER) diffraction meter. Microanalysis of specimen was performed with Neophot-2 metallographic microscope with 200x, 500x and 1000x magnification [4,5,6].

EXPERIMENTAL PART

To try out the new alloy technology involving the addition of CrB₂, the team used self-fluxing surfacing powder alloy PG-Z40 with 50-160 μm grading fraction and the following chemical composition: Fe=38%; Cr=15,0%; Si=3,1%; B=2,9%; Ni=33%; Cu=4%; V=4,0%.

Addition alloy CrB₂ with the following chemical composition: Ti=0,08%; Fe=0,47%; Ni=0,52%; B=15%; Cr=83,43% was used as a hardening additive. CrB₂ hardener was granulated for 5 minutes in a laboratory planetary granulator in alcohol environment with the use of grinding media (5-8 bearing balls with 1:3 powder mix / balls weight ratio). After the granulation average size of CrB₂ particles was 10-30 μm.

2 types of powder mix were prepared for the experiment.

The first type of powder mix was the new surfacing alloy PG-Z40, and the second type was PG-Z40 with CrB₂ hardener. The powder mix was a mechanical mixture of the new PG-Z40 alloy initial state components and CrB₂ hardener subjected to

mechanical alloying in an attritor. Mechanochemical activation of the powder mix was also carried out in attritor. Prior to the mechanochemical activation, 1% of zinc stearate was added to the powder mix to prevent mix material sticking to the balls and the inner surface of the drum. The mix was treated in attritor with the following production conditions:

- Mixer rotation rate – 340 rpm;
- Ball diameter – 5 mm;
- Powder mix weight / ball weight ratio – 1/18;
- Process duration – 2 hours.

Mechanical alloying produced composite powder materials with particle size 10-20 μm [5-8]. To produce glomed powder surfacing material, the initial powder material obtained in the attritor was mixed with an organic bond of 2-3% alcohol solution of phenolic varnish FL-98 with subsequent sintering in a LH15/12 batch furnace at the temperature of 700°C [7,8].

After the sintering, the glomed material was granulated and sieved through a set of sieves to sort out 50-160 μm size particles. The testing of experimental boride-hardened powders for gas powder surfacing was prepared in accordance with GOST 21448-75 «Surfacing alloy powders» taking into account existing methods [9].

Surfacing was performed by oxypropane torch with No.5 tip manufactured under the specification No. 200 of Kaz. SSR 210-84, under the patent №1276 issued by the Republic of Kazakhstan. Compressed gases were used during surfacing: oxygen under GOST 5583-78 and propane under GOST 20448-80. The reference specimen was made of 45 steel under GOST 1050-88, and had dimensions of 30x45x11 (mm). Gas powder surfacing was carried out by the application of 1,2-1,5 mm thick surfacing layer under the following conditions [10]:

- Oxygen pressure at the torch inlet - 9,0 kg/cm²;
- Propane pressure at the torch inlet - 1,2 kg/cm²;
- Oxygen expenditure – 750 l/h;
- Propane expenditure – 700 l/h;
- New surfacing alloy expenditure – 50 g/min;
- Surfacing layer thickness – 2-3 mm.

The surface of the surfacing layer was treated with an abrasive tool made of green silicon carbide. The coating is free of pinholes and slag inclusions.

RESEARCH RESULTS AND DISCUSSION

Fig.1 contains photomicrographs of the produced surfacing powder demonstrating that the particles are globe- and oval-shaped and their size ranges from 50 to 160 μm.

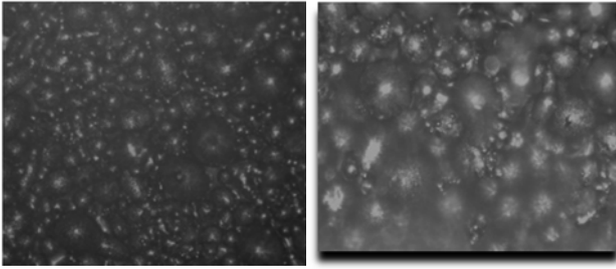


Fig.1. Photomicrographs of the powder with x200 and x500 magnification

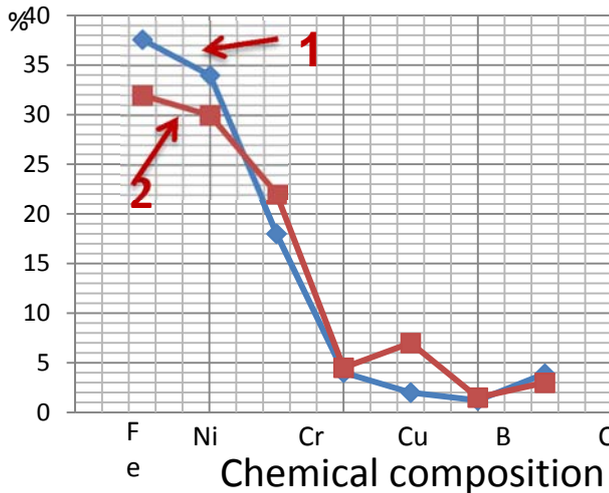


Fig.2. Chemical composition of the surfacing powder

- 1- Powder alloy PG-Z40 – 100%;
- 2- Powder alloy PG-Z40 90% + 10% CrB₂

Chemical composition (Fig.2), powder and composite coating structure was studied with a spectrometer. The used spectrometer had no function of boron, carbon and silicon identification, therefore the initial data on their content in the studied alloys

was put in the graph. The content of other elements in the alloy corresponds to their content in the initial powder. The effect of CrB₂ additive on the changes in the chemical composition of the PG-Z40-based alloy is shown on the graph.

X-ray structure analysis (Fig.3) showed that the alloy's structure is a composition including relatively strong and plastic iron nickel matrix and strengthening phases in the form of carbides and borides such as FeB; Cr₂B; Cr₂B₂; Cr₃C₂; Cr₅B₃; Cr₇C₃; Cr₂₃C₆; Fe₃C; Cr₇BC₄; Fe₃C, silicon nickelide Ni₂Si₅. The produced material is an iron nickel low alloy with a typical two-phase structure. The 12,1% increase of chromium content in the two-phase Fe-Cr structure of the PG-Z40+10% CrB₂ alloy in comparison with the similar PG-Z40 structure should also be noted.

Microhardness on Vickers scale was measured separately for the two alloys and the diffusion zone with the results showing hardness value on the steel baseplate in the diffusion zone and on the surface of the studied glomed powder alloy. The obtained microhardness values for the surfaced coating in the fusion zone of the studied specimen are shown in the Tab.1.

It has to be noted that microhardness increases in the direction towards the surfaced coating of the studied sample.

Coating microhardness values are uniform, there are no apparent hardness variations. The value range from 400 to 500 MPa, implying uniform distribution of carbides and borides, i.e. the structure is homogeneous throughout the entire area of the surfacing coating (Fig.4).

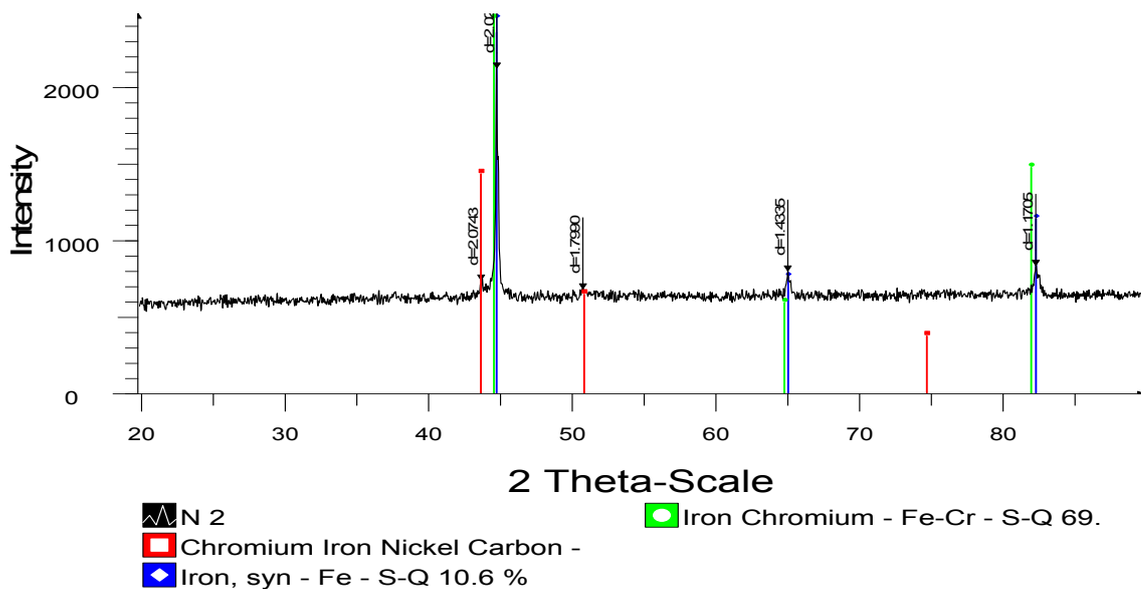


Fig.3. The results of X-ray structure analysis for PG-Z40+10% CrB₂

Table 1.Microhardness measurement results for the studied sample.

Impressions	Vickers hardness, HV 10 kgf		
	Steel baseplate	Diffusion zone	Sample material, 90% PG-Z40+10% CrB ₂
1	382.72	381.40	456.72
2	362.75	426.57	470.63
3	375.41	428.62	530.29
4	359.34	443.42	485.18
Average	HV = 370.34	HV =420.00	HV =546.96

The coating microhardness was measured on the polished specimen №1, №2 having 2 mm thickness in 4-7 measurements of the diagonal of impression.

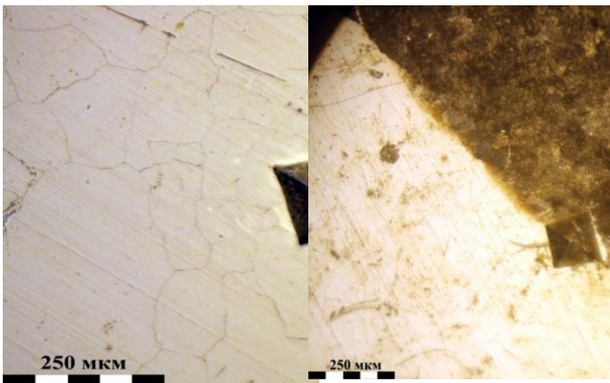


Fig.4.The size of an impression on the polished specimen of the samples: a) PG-Z40; b) PG-Z40+10% CrB₂

The obtained microhardness values for the surfaced coating of the sample are shown in the Tab.2.

Table 2.Microhardness measurement results for the polished specimen of the sample №1 surfaced with PG-Z40 alloy

Impressions	Vickers hardness, HV 10 kgf	
1	482.71	563.64
2	519.13	546.15
3	487.67	542.09
4	492.71	547.96
Average	HV=495.55	HV=549.96

According to microhardness measurement results, the average microhardness value of the sample surfaced with the boride-hardened alloy increases by 54,41 HV in comparison with the initial alloy obtained with the surfacing with PG-Z40 material.

The study of the fusion zone of the new boride-hardened alloy was carried out with the use of metallographic microscope with x200 and x500 magnification. The polished specimen was prepared using a sample surfaced with glomed powder alloy of

the following composition: 90% PG-Z40+10% CrB₂. Fig.5 shows the boundaries of the surfaced alloy's diffusion zone. The formed diffusion zone layers seem to be the result of peritectic reaction, in accordance with the system state diagram: Fe-Ni-Cr-B.

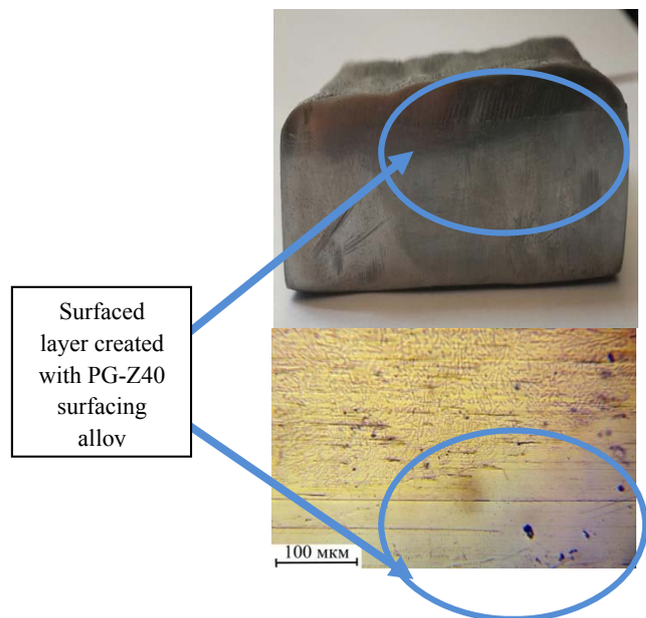


Fig.5. Microstructure of the surfaced metal (PG-Z40+10% CrB₂) with base metal, x500

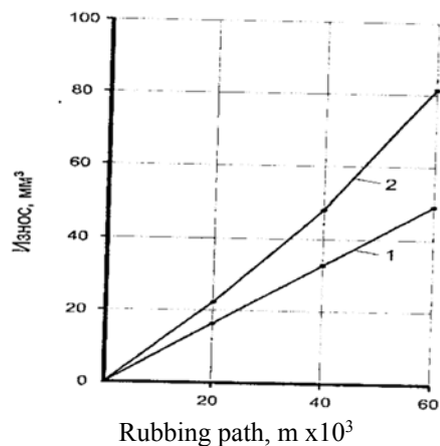


Fig.6. The new alloy and steel wear rates: 1-The new PG-Z40+10% CrB₂ alloy 2-45 steel, 55 HRC hardness

Fig.6 shows the results of wear assessment for the samples with surfaced coating hardened to 55 HRC compared to 20GL steel dressers. We can assume that wear resistance and durability of components of friction units is increased after their reconditioning by PG-Z40 +10% CrB₂ alloy surfacing.

Metallographic analysis was conducted for the sample No.1 surfaced with PG-Z40 alloy and for the sample No.2 surfaced with PG-Z40+10% CrB₂ alloy. Microanalysis was performed with the use of Neophot-32 metallographic microscope with $\times 200$ and $\times 500$ magnification.

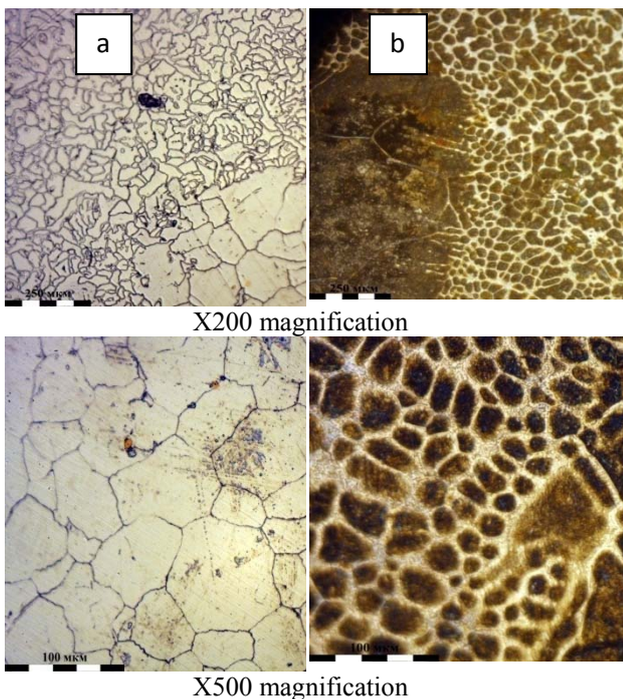


Fig.7. Microstructure of the layer surfaced with PG-Z40 alloy (a) and PG-Z40+10% CrB₂ alloy

Fig.7 shows homogeneous polyhedral, equiaxial block structure with a coarse-grain to fine-grain structure transition zone and localized solid copper solution particles. Chains of multiple proeutectoid constituent precipitations of two types (dark, and sometimes light), which seem to be iron and chromium carbides, can be observed along the grains' boundaries.

Fig.7 also shows large crystals composed of ultrafine-grain sub-crystals in the microstructure of the alloy. Polyhedral ferrite crystals – solid carbon solution in α -iron and perlite – possibly represent eutectoid mix of ferrite and iron and chromium carbide.

The phases mainly formed inside the crystals and along their boundaries in some places are as big as 1 μm and form block structure. From the obtained data we can conclude that the developed boride-hardened

surfacing powder PG-Z40+10% CrB₂ has increased hardness and better properties compared to PG-Z40 surfacing powder [11].

CONCLUSIONS

1. The research helped to determine an optimal composition for CrB₂ hardener added to the new PG-Z40 surfacing alloy. It has been determined that in order to achieve 450-600 HV hardness of surfaced metal, the hardener has to be added in the amount up to 10% of total weight.

2. X-ray structure analysis showed that after mechanical alloying and melting of the compound, the composite surfacing alloy PG-Z40 with the addition of 10% of chromium boron appears as an iron nickel matrix with evenly distributed iron, chromium and nickel borides and carbides, and this observation is confirmed by the surfaced metal microhardness increase from 495,55 to 549,96 HV.

3. Increased hardness coatings can be created on the basis of the self-fluxing PG-Z40 alloy by adding chromium borides to the alloy prior to gas powder surfacing. Chromium borides are genetically bound by the composition of the surfacing material. The obtained structure ensures maximal wear-resistance of the coating and is characterized by even distribution of hardening crystal throughout the entire section achieved by the coating layer surface melting at the temperature of 1000-1100°C. Inadequate temperature during melting does not dissolve the added borides, and it results in non-homogeneity of the surface microstructure.

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