Soft magnetic polymer-metal composites consisting of nanostructural Fe-basic powders

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ABSTRACT

Purpose: The paper presents and reviews the research results of soft magnetic composites consisting nanocrystalline powders obtained by soaking and high energetic milling of amorphous ribbons of metallic glasses Fe79Si3B13 and Fe73.5Cu13.5Nb13.5B9.

Design/methodology/approach: Amorphous Fe79Si3B13 and Fe73.5Cu13.5Nb13.5B9 ribbons were milled in a high energy ball mill (8000 SPEX CertiPrep Mixer/Mill) with a ball-to-sample weight ratio of 5:1. The obtained metallic powders were sieved to a particle mean diameter 200–500 μm, 75–200 μm and 25–75 μm, and then annealed in an argon atmosphere to generate the nanocrystalline state. The powders particles were mixed and consolidated with polymer to obtain composites in the form toroidal cores. Observations of the structure of powders and composites were made on the Optron DSM-940 scanning electron microscope and electron transmitting microscope JEOL JEM 200CX and X-ray analysis. The X-ray tests were realized with the use of the XRD 7 SEIFERT-FPM diffractometer.

Findings: The analysis of the magnetic properties test results of the powders obtained in the high-energy ball of milling process, and the composites manufactured from these powders proved that the process causes significant decrease in the magnetic properties in relation to ribbons. The structure and magnetic properties of this material may be improved by means of a proper choice of parameters of this process as well as the final thermal treatment and first of all by decrease of demagnetization effect.

Research limitations/implications: For the powders, further magnetic, structure and composition examinations are planned.

Practical implications: The amorphous and nanocrystalline Fe79Si3B13 and Fe73.5Cu13.5Nb13.5B9 powders obtained by high-energy ball milling of metallic glasses feature an alternative to solid alloys and make it possible to obtain the ferromagnetic nanocomposites, whose shape, dimensions and magnetic properties can be freely formed in the determinate range.

Originality/value: The paper presents influence of chemical composition of selected Fe-based metallic glasses, annealing temperature and parameters of the high-energy ball milling process on structure and magnetic properties of soft magnetic powders materials obtained in this technique. This paper review also possibility of magnetic properties forming for metal-polymer nanocrystalline composites. Results and a discussion of the influence of high energy mechanical milling process on particle size and their distribution and annealing temperature of powders as well as structure and magnetic properties of investigated samples is presented. According to achieved results it has been attempted to describe the possibilities of improvement the soft magnetic properties of obtained Fe-based nanocrystalline powder materials and composites from them manufactured.

Keywords: Nanomaterials; Powders; Heat treatment; Magnetic properties; Magnetic composites
1. Introduction

In recent years, the Fe-based nanostructural (nanocrystalline) alloys has been intensively studied for its excellent soft magnetic properties [1÷10]. These nanostructural materials are prepared from amorphous ribbons approximately 20-50 µm thick produced by melt spinning method and next annealing or in form of powders by ball milling of metallic glasses ribbons or by mechanical alloying [1,11÷13]. The magnetic properties of the metallic glass ribbons and nanostructural materials depend on the chemical composition, production procedure and on the condition of the heat treatment (time and temperature) or parameters of ball milling [14÷16]. After annealing the ribbons of metallic glasses at above crystallization temperature for 1 hour in argon protecting atmosphere they become nanocrystalline. But due to process of rapidly solidification the shape of these alloys is limited practically to ribbons and wires (except bulk metallic glasses). The ribbon and wire geometries are very satisfactory for a range of applications, such as sensors and pulse generators, but are not the best for those in which large mass or volume of soft magnetic materials or complexly shaped soft magnetic part are required. Positive outcome of the amorphous – nanocrystalline transformation is the improvement of soft magnetic properties like very low coercitivity of less than 1 A/m, low AC losses and high magnetic saturation compared with amorphous materials. On the other hand, the ribbons of metallic glasses become very brittle, restricting its possible applications. Second disadvantage for many applications is the small thickness of ribbons and shapes of cores which are generally limited to toroidaly wound or stacked types [1÷10].

Obtaining powder nanocrystalline materials directly by grinding the metallic glasses makes it possible to carry out research on obtaining the ferromagnetic nanocomposites whose shape and dimensions may be formed freely [17÷25].

The composite with the polymer matrix reinforced with the amorphous or nanocrystalline particles obtained in the process of the high energy milling of the amorphous strips or mechanical alloying features the alternative for those materials.

Investigation of composite materials obtained by consolidation of the metallic powders using various methods is carried out in many research centers in the world [17÷25].

This solution encounters many difficulties, as particles with the length to diameter ratio of 1:1 (or close to it) run the significant risk of the demagnetising phenomenon.

In the case of soft magnetic a powder material, during the magnetisation of the short cylindrical sample, the saturation is achieved at the magnetic field intensity much greater than in the case when the close ring is used. It is caused by the fact that the intensity of the magnetic field \( \mu^0 \cdot H \) in the short sample is lower than the magnetic field \( \mu^0 \cdot H \) in the whole coil [23].

There is a relation between the magnetization \( J \) and the field intensity. When the partial demagnetization caused by the shape of the sample is taken into account, the relation may be presented in the following way:

\[
J = \mu_0 \cdot H \cdot \frac{k}{1 + \kappa N}
\]

where:
- \( N \) - demagnetization coefficient
- \( k \) - magnetic susceptibility
- \( \mu_0 \) - induction constant (magnetic permeability of vacuum).

The \( N \) coefficient in the case of, for example, cylindrical samples depends mainly on the relation of sample length \( l \) to its diameter \( D_p \) [23].

\[
P = \frac{l}{D_p}
\]

The shorter and thicker the cylindrical rod is, the smaller value of the \( p \) parameter and at the same time the higher value of the \( N \) parameter is. The demagnetization effect of sample ends is presented in Fig.1.

![Fig. 1. The effect of demagnetisation of flat sample ends [23]](image)

In addition, in case of the very small particles, below 1 µm the superparamagnetism phenomenon may occur. Both these phenomena deteriorate the soft magnetic properties of the obtained composites. Nevertheless, apart from the conventional heat treatment methods and heat treatment in the magnetic field, regulation is possible (in a certain range) of the magnetic and mechanical properties of the composite by selection and segregation of the powder grains according to their sizes and by selection of the mass portion of the polymer matrix.

The ball milling techniques of metallic glasses ribbons is often used to produce nanocrystalline soft magnetic material which is in powder form, and therefore is suitable for consolidation in a variety of shapes [14÷17]. Different consolidation techniques have been reported recently [16÷21]. The proposed procedures like sintering, hot isostating pressing, warm compaction, explosive compaction, shock – wave compaction and static high – pressure compaction with pressures up to 5 GPa are costly and complex. Processing of bulk materials from nanocrystalline powder (e.g. sintering) is very difficult due to the metastable state of nanocrystals. Annealing at high temperatures causes grain growth and soft magnetic properties decrease. That is why the most popular consolidation technique is mixing metallic powders with mineral or polymer binder [19÷25].

The optimization of the magnetic properties of nanocrystalline powder composites can be achieved by adjusting uniform packing density, the powder size, the volume binder rate, the nature of the binder and ideal orientation of powders particles in the cores.

The aim of this study is the preparation of toroidal shape nanostructural composite cores from high energy ball milled \( \text{Fe}_{73.5}\text{Cu}_{1}\text{Nb}_{5}\text{Si}_{13.5}\text{B}_{2} \) powders and to investigate the powder particle size and content of polymer binder response of the soft
magnetic properties. The results of scanning electron microscopy (SEM), transmission electron microscopy (TEM) and X-ray diffraction (XRD) of the obtained powders is also presented.

The amorphous and nanocrystalline materials arouse interest of scientists in research centers all over the world in the last three decades. This is connected with the soft magnetic properties characteristic for these materials and with the possibility of using them in the electronic and electrical industry [1÷4].

The metallic amorphous and nanocrystalline materials obtained by crystallization of the metallic glasses are available in the form of thin ribbons only, and their magnetic properties can be controlled by heat- or thermo-magnetic treatment. These limitations are complicated in addition by the feasibility of manufacturing only toroidal cores from the ribbons.

Obtaining the nanocrystalline metal powders by milling of metallic glasses features an alternative to solid materials like amorphous and nanocrystalline ribbons and makes it possible to obtain the ferromagnetic nanocomposites, whose shape and dimensions can be freely formed using various consolidation methods [14÷25].

2. Materials and methods

Amorphous Fe₇₃Si₉B₁₇, Fe₇₅Cu₁Nb₃Si₁₃B₉, metallic glasses ribbons (fig.2,3), were prepared by the melt spinning method. The width and thickness of these ribbons were 9-10.2 mm and 28-29 µm, respectively. A Thermolyne F6020C resistance muffle furnace was used for isothermal soaking of the amorphous ribbons and the powder. Soaking was carried out at temperature range 373-873 K for 1 hour in an argon protective atmosphere.

The annealed ribbons of metallic glasses were cut into small pieces of about 5 mm transverse dimension. These particles were sealed in a cylindrical stainless steel container under an Ar atmosphere. The ball milling process was performed in a high energy ball mill (8000 SPEX CertiPrep Mixer/Mill) with a ball-to-sample weight ratio of 5:1. The milling process reduced the starting material pieces (metallic glasses) to powder form. The metallic powder particles were sieved to obtain different sizes – small with average particles diameter between 25 to 75 µm, medium particles 300÷500 µm and the large ones between 750÷1200 µm. Obtained powders were mixed then with different contents of binder and condensed on mechanical shaker. As a binder, a silicone polymer Dublisil 20 from Dreve-Dentamid GMBH and other polymer materials was used.

The powder particles to silicon polymer weight ratios were 6:1, 5:1, 4:1, 3:1, 2:1 and 1:1 respectively. The toroidal cores so obtained had its external diameter of 34 mm, internal diameter of 28 mm and height of 8 mm. Fig. 1 shows the consolidated samples of toroidal cores.

X-ray examination of metallic ribbons and obtained powders was made using a XRD-7 (SEIFERT-FPM) diffractometer with a Cu anode, producing radiation of λ = 0.17902 nm wave length. The X-ray tube was supplied with the current I = 40 mA under the voltage of U = 35 kV. Diffraction examinations were performed within the range of angles from 20 to 20 to 120 degrees. The measurement step was of 0.2° in length whilst the pulse counting time was 10 s.

The size of crystallites was determined using the Scherrer formula [1].

The morphology of the powder samples was observed by scanning electron microscopy OPTON DSM 940.

Examination of powders and composite cores magnetic properties (hysteresis loops) were carried out on the Lake Shore Cryotronics VSM vibration sample magnetometer with a maximum applied magnetic field of 800 kA/m. The permeability of the powders and composites cores was made on the Ferrometr device.
3. Structure and properties of powders

Both milling method of metallic glasses ribbons: directly as quenched state and after annealing, conducted to getting powders with nanocrystalline structure (nanostructural powders) with different combination of phase composition; amorphous, Fe-α or α - Fe(Si) and borides, with different size of powders particles and different size of nanocrystals in amorphous matrix, depend from time of milling and time and temperature of soaking of metallic glasses ribbons.

The morphology of the sieved powders obtained by milling amorphous ribbons examined by scanning electron microscopy (SEM) showed significant differences in the shape and size of powder particles in relation to the milling time. It is important because influence for the demagnetization effect and magnetic properties of powders. Fig. 4 A,B,C shows SEM images of Fe$_{78}$Si$_9$B$_{13}$ powder particles sieved to obtain different sizes – small with average particles diameter between 25 to 75 µm, medium particles 75÷200 µm and the large ones between 200÷500 µm Fig.4 D,E,F shows SEM images of Fe$_{71.5}$Cu$_1$Nb$_1$Si$_{13.5}$B$_9$ powder particles sieved to obtain different sizes – small with average particles diameter between 25 to 75 µm, medium particles 300÷500 µm and the large ones between 750÷1200 µm.

It was found that larger powder particles had shape like flakes as well as plates with sharp edges. Their thickness was much smaller than dimensions in two remaining directions. Shape of medium particles was also similar to the shape of flakes. Powders with average particles diameter between 25 to 75 µm had very different shape – irregular flakes, flat with round edges and perfect spherical shape, suitable for forming consolidated powder cores with high density. It was also found that some of the powder particles were agglomerated.

The average particle size for the Fe$_{78}$Si$_9$B$_{13}$ powder in the particle size distribution curve was for small particles (25÷75 µm) about d = 48 µm, for medium particles (75÷200 µm) about d = 127 µm and for the large ones (200÷500 µm) about d = 326 µm. For example the average particle size for the Fe$_{71.5}$Cu$_1$Nb$_1$Si$_{13.5}$B$_9$ powder in the particle size distribution curve was for small particles (25÷75 µm) about d = 58 µm, for medium particles (300÷500 µm) about d = 385 µm and for the large ones (750÷1200 µm) about d = 980 µm (Fig.5).

To obtain information about the crystalline phases an X-ray diffraction study (Fig.6) was carried out for the base sample (Fe$_{78}$Si$_9$B$_{13}$ metallic glass) for ball milled ribbon and for obtained powders annealed in Ar for 1h at temperature of 773 K.

It was found for the rapidly quenched ribbon the occurrence of a wide diffraction spectrum within the range of 2θ = 45 ÷ 58°, being that phenomenon characteristic for an amorphous structure of iron alloys (A in Fig. 6). Fig 6 (B) shows the X-ray diffraction pattern for amorphous Fe$_{78}$Si$_9$B$_{13}$ powders. Fig 6 (C) shows the X-ray diffraction pattern for nanocrystalline Fe$_{78}$Si$_9$B$_{13}$ powders annealed in Ar for 1h at temperature of 773 K. There was a sharp peak with high intensity observed in the curve for the angle 2θ = 52.8° being the characteristic angle for the α-Fe(Si) (110) plane reflection. Also (200) and (211) diffraction peaks of α-Fe(Si) were noticed.

Similar X-ray investigations performed for Fe$_{78}$Si$_9$B$_{13}$ and Fe$_{71.5}$Cu$_1$Nb$_1$Si$_{13.5}$B$_9$ powders manufactured by high energetic milling of metallic glasses ribbons showed similar results as above mentioned what very good illustrate Fig.7,8.

After 50 hours of Fe$_{78}$Si$_9$B$_{13}$ metallic glasses ribbon the existence of amorphous and α-Fe(Si) phases, which share changing with the time of milling. After 100 hours of milling probably advance process of amorphous phase crystallization occurring. Opposite is for Fe$_{71.5}$Cu$_1$Nb$_1$Si$_{13.5}$B$_9$ metallic glasses ribbon where after 100 hours milling very small changes are observed, which is evidence that structure is further amorphous.

The average crystallite size in the obtained powder was determined from the half maximum intensity breadth of the dominant peak of the X-ray diffraction pattern, using the Scherrer formula and amounts about 20-33 nm.
After soaking the Fe$_{73.5}$Cu$_1$Nb$_3$Si$_{13.5}$B$_9$ material at 823 K for 1 hour, a high-energy grinding was carried out for 1, 2, 3, 4 and 5 hours respectively. A nanocrystalline powder was obtained as a result of the combination of thermal nanocrystalline and milling processes. The size of crystallites so obtained in the powder were function of time milling, and were smaller for longer time of milling (Fig.9), and share of crystalline and amorphous phases was also changing.

It was also found, that in high-energy milling of the amorphous Fe$_{78}$Si$_{13}$B$_9$ alloy strip soaked at 450°C for 1 hour, the intensity of the Fe-(110) diffraction lines decreases whilst small peaks from the Fe$_3$B phases appeared (Fig. 10).

Generally from this experiment we can conclude that is possible to fabricate nanocrystalline powders from metallic glasses with different size of powder particles and different structure of powders by the use of a combined thermal nanocrystallization and high-energy milling, and this way we can minimize production time of determine powders.
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Fig. 5. Average diameter of powder particles vs. milling time of metallic glasses ribbons

Fig. 6. X-ray diffraction patterns of Fe$_{78}$Si$_{13}$B$_{9}$ samples: as-quenched ribbon; (B) powders obtained from ribbon; (C) powders annealed at 773 K for 1 hour

After soaking the Fe$_{73.5}$Cu$_{1}$Nb$_{3}$Si$_{13.5}$B$_{9}$ material at 823 K for 1 hour, a high-energy grinding was carried out for 1, 2, 3, 4 and 5 hours respectively. A nanocrystalline powder was obtained as a result of the combination of thermal nanocrystalline and milling processes. The size of crystallites so obtained in the powder were function of time milling, and were smaller for longer time of milling (Fig. 9), and share of crystalline and amorphous phases was also changing.

It was also found, that in high-energy milling of the amorphous Fe$_{78}$Si$_{13}$B$_{9}$ alloy strip soaked at 450°C for 1 hour, the intensity of the Fe-$\beta$ diffraction lines decreases whilst small peaks from the Fe$_3$B and Fe$_2$B phases appeared (Fig. 10).

Generally from this experiment we can conclude that is possible to fabricate nanocrystalline powders from metallic glasses with different size of powder particles and different structure of powders by the use of a combined thermal nanocrystallization and high-energy milling, and this way we can minimize production time of determine powders.

Fig. 7. A set of x-ray diffractions for powders fabricated from the Fe$_{78}$Si$_{13}$B$_{9}$ metallic glasses strip for different time of milling

Fig. 8. A set of x-ray diffractions for powders fabricated form the Fe$_{73.5}$Cu$_{1}$Nb$_{3}$Si$_{13.5}$B$_{9}$ metallic glasses strip for different time of milling

Fig. 9. X-ray diffraction patterns of Fe$_{73.5}$Cu$_{1}$Nb$_{3}$Si$_{13.5}$B$_{9}$ alloy the annealed sample at 823 K and milled for 1, 2, 3, 4, 5 hours

Fig. 10. A set of diffractograms for powders fabricated from an amorphous Fe$_{78}$Si$_{13}$B$_{9}$ alloy strip soaked at 450°C for 1 hour

Fig. 11. Structure of a Fe$_{78}$Si$_{13}$B$_{9}$ alloy soaked at 450 °C/1 hour; TEM micrograph.

Fig 12. Transmission electron microscopy bright field image and selected area electron diffraction pattern of Fe$_{78}$Si$_{13}$B$_{9}$ powders annealed at 773 K for 1 hour.

Fig 13. Transmission electron microscopy bright field image and selected area electron diffraction pattern of powders obtained from Fe$_{73.5}$Cu$_{1}$Nb$_{3}$Si$_{13.5}$B$_{9}$ ribbon annealed at 823 K.

Fig.10. A set of diffractograms for powders fabricated from an amorphous Fe$_{78}$Si$_{13}$B$_{9}$ alloy strip soaked at 450°C for 1 hour

Fig.11 shows the transmission electron microscopy (TEM) microstructure image and the corresponding selected area electron diffraction (SAED) pattern of the ball milled Fe$_{78}$Si$_{13}$B$_{9}$ sample annealed in Ar for 1h at temperature of 773 K. The TEM image demonstrated nanocrystallits with a size around 20 nm embedded in an amorphous matrix. It was consistent with SAED pattern, which exhibited diffraction rings for the (110), (200) and (211) of $\alpha$-Fe($\beta$) phase and a diffuse amorphous ring (Fig.12). The phases identified by TEM were a good agreement with XRD analysis results.

Transmission electron microscopy (TEM) observation of the as-quenched Fe$_{73.5}$Cu$_{1}$Nb$_{3}$Si$_{13.5}$B$_{9}$ ribbon confirmed that the microstructure of the sample is very uniform. The bright field image shows a homogeneous material with no crystalline grains. Also, there is no trace of crystalline reflections in the selected area electron diffraction (SAED) pattern. These factors indicated that the as-quenched ribbon is fully amorphous.

Fig.13 shows the TEM microstructure image and the corresponding selected area electron diffraction (SAED) pattern of the ball milled Fe$_{73.5}$Cu$_{1}$Nb$_{3}$Si$_{13.5}$B$_{9}$ sample (previously annealed in Ar for 1h at temperature of 823 K). The phases identified by TEM were a good agreement with XRD analysis results.
The saturation induction $B_s$, remanence $B_r$ and coercive field $H_c$ were determined from magnetic hysteresis loops of powders measured in a VSM for powders Fe$_{78}$Si$_{13.5}$ and Fe$_{73.5}$Cu$_9$Nb$_3$Si$_{13.5}$B$_9$ with different structure produced by different methods and at different parameters (for example Fig.14,15).

Very interesting from the point of view demagnetization effect is examination of the magnetic properties of the both alloys powders as a function of particles size. For the Fe$_{78}$Si$_{13.5}$ powders were found that the larger powder particles (200 ÷ 500 µm) exhibits better soft magnetic properties than the small ones i.e. the coercive field $H_c = 1341$ [A/m], $B_r = 1.06$ [T] and $B_s = 0.068$ [T]. Decreasing the particles size to 75 ÷ 200 µm decreased the $B_r = 0.98$ [T] and $B_s = 0.061$ [T]. Also increasing of coercive field up to 1920 [A/m] was observed. For small particle size (25 ÷ 75 µm) powder sample demonstrated the value of $B_r = 0.72$ [T] and the remanence $B_s = 0.048$ [T], whilst $H_c = 3215$ [A/m].

The permeability value of large and medium powder particles (measured at magnetic field $H = 3$ [kA/m] and at frequency $f = 50$ Hz) are $\mu = 39$ and $\mu = 18$ respectively. Annealing powders particles at 773 K produces the effect of a considerable increase of their magnetic properties. For large particles the value of coercive field was equal to 237 [A/m], $B_r = 1.22$ [T], $B_s = 0.09$ [T] and the permeability exhibited value of 106 (Fig.16).

Similar investigation of the magnetic properties of the Fe$_{73.5}$Cu$_9$Nb$_3$Si$_{13.5}$B$_9$ alloy powders with different particles size also showed that the larger powder particles exhibits better soft magnetic properties than the small ones i.e. the coercive field $H_c = 13.75$ [A/m], the saturation induction $B_r = 1.21$ [T] and the remanence $B_s = 0.007$ [T]. Decreasing the particles size to 300 ÷ 500 µm decreased the saturation induction down to $B_r = 1.06$ [T] and $B_s = 0.006$ [T]. Also increasing of coercive field up to 115.3 [A/m] was observed (Fig.17). For small particle size powder sample demonstrated the value of $B_r = 0.64$ [T] and the remanence $B_s = 0.0038$ [T], whilst $H_c = 163.2$ [A/m].

The permeability value of large and medium powder particles (measured at magnetic field $H = 3$ [kA/m] and at frequency $f = 50$ Hz) are $\mu = 175$ and $\mu = 143$ respectively.

For both alloys powders the magnetic permeability measurements confirmed better magnetic properties of powder particles, which had shape like flakes.

![Fig. 14. Hysteresis loops for Fe$_{78}$Si$_{13.5}$ powder particles (200 ÷ 500 µm) size, in as milled state and after annealed at 773K for 1 hour](Image)

![Fig. 15. Hysteresis loops for Fe$_{73.5}$Cu$_9$Nb$_3$Si$_{13.5}$B$_9$ alloy powders with different particles size](Image)

![Fig. 16. The influence of the soaking temperature on the coercive force $H_c$ and the maximum magnetic permeability $\mu_{max}$ of a Fe$_{78}$Si$_{13.5}$B$_9$ alloy powders.](Image)

![Fig. 17. The influence of the milling time on the coercive force $H_c$ of the Fe$_{73.5}$Cu$_9$Nb$_3$Si$_{13.5}$B$_9$ alloy powders.](Image)
4. Composites research results and discussion

The investigations of magnetic properties performed for metal-polymer composites, consisting above described powders showed very strong relation of powder particle size and share of ferromagnetic phase on magnetic properties of composites. The structure and level of powders magnetic properties influence weaker for magnetic properties of composites.

The results of examination of the magnetic properties of the nanocomposite cores with different powder-to-silicone polymer ratio are shown in Fig. 18-23.

The coercive field (Fig.18,19) and the saturation induction (Fig. 20,21) of powder composite cores in dependence on the powder particles content (wt %) are compared.

The nanocrystalline cores with powder particles between 750÷1200 µm exhibited high value of saturation induction and coercive field was comparable for the powders obtained from ball milled ribbon previously annealed in argon protecting atmosphere for 1h at temperature of 823 K. The results of experimental studies demonstrated a correlation between the growing powder-to-silicon polymer ratio and the improvement of the magnetic properties of the composite.

The composite core with a powder particle content of 85.7 wt% exhibit Hc=83.2 [A/m], B0=1.09 [T] and B0=0.0058 [T]. Further decrease powder-to-silicon polymer ratio leads to deterioration of the magnetic properties, i.e. the coercive field goes considerably high at small values of both the remanence and the saturation magnetic induction: for example, for a composite core with a polymer content of 50 wt%, B0=0.0049 [T], Bc=0.95 [T] whilst the coercive field 214.3 [A/m].

By decreasing the particles size fraction to 300-500 µm the coercive field of composite core with powder particle content of 75 wt% is increased at low values of both the remanence and the saturation magnetic induction. For medium particle size, composite cores with different content of polymer binder showed Hc in the range 163-267 [A/m], the values of Bc between 0.88-0.97 [T] and the remanence B0 in the range 0.0048-0.0054 [T].

Toroidal cores filled with small powder particles (25-75 µm) demonstrated worse magnetic properties than the other samples. At 85.7 wt% powder the coercive field Hc was equal 238.1 [A/m], Bc=0.62 [T] and B0=0.0032 [T].

Decreasing powder content to 50 wt% decreased the value of Bc to 0.51 [T] and B0 to 0.0024 [T], whilst Hc=311.4 [A/m].

The magnetic permeability measurements confirmed better magnetic properties of nanocomposite cores with powder particles between 750÷1200 µm (Fig. 23). These data represent the highest reported permeability (measured at magnetic field H=3 [kA/m] and at frequency f=50 Hz) μ=123 for core with Fe73,5Cu13Nb13.5Si13B9 powder particle content of 85.7 wt%.

The medium powder particles (300-500 µm) in the composite cores showed permeability between 64 and 106 depending on the powder-to-silicon polymer ratio. It was also found that for powder particles size between 25-75 µm, composite cores with different content of polymer binder showed permeability in the range 20-48 (measured at magnetic field H=3 [kA/m] and at frequency f=50 Hz).

Magnetic properties very strong depend from the demagnetization effect (powder grain size and shape) then from structure of powders particles, therefore impossible is to rich for polymer-metal composites similar magnetic properties as for winded cores of metallic glasses or nanostructural ribbons. However we can have other advantages from polymer-metal magnetic composites; possibilities of easy forming different shapes of magnetic cores, and much lower the active power losses (Fig.24,25).

5. Conclusions

The experimental studies showed that a combined process involving the thermal nanocrystallization and high-energy ball milling applied to an Fe78Si5B13 and Fe73.5Cu13Nb13.5Si13B9 alloys provided a solution to obtain a metallic powder containing α-Fe(Si) crystallites with a size around 20 nm embedded in an amorphous matrix. This two phase structure, present in the powder obtained by ball milling of nanocrystalline ribbons, was the optimal structure from magnetic properties point of view.

It was found that size and shape of the powder particles were responsible for soft magnetic properties of powder cores. Decreasing size of powder particles led to magnetic hardening of nanocrystalline Fe78Si5B13 and Fe73.5Cu13Nb13.5Si13B9 alloys. The main reason for the worse magnetic properties of small particles was high demagnetization factor. Additionally, structural defects induced during ball milling process of small particles also caused deterioration of magnetic properties.

The control of the soft magnetic properties of composite cores consolidated from nanocrystalline powders and silicone polymer binder can be achieved by adjusting powder particle size and powder-to-silicon polymer ratio.

It was found that composite cores fabricated from silicon polymer-solidified Fe78Si5B13 and Fe73.5Cu13Nb13.5Si13B9 nanocrystalline powders exhibits magnetic properties that strengthen with the growing metallic powder-to-silicon weight or volume ratio.

The best results (Hc=83.2 [A/m], Bc=1.09 [T], B0=0.0058 [T] and μ=123 measured at magnetic field H=3 [kA/m] and at frequency f=50 Hz) were obtained for the composite core filled with powder particles between 750÷1200 µm and with a powder particle content of 85.7 wt%. These results are similar as for composites investigated by other authors for polymer-metal composites manufactured without pressing (Fig.26).

Magnetic properties (particularly magnetic permeability) very weak depend from initial magnetic permeability of metallic glasses ribbons used to this investigation.
Fig. 18. Influence of the content of Fe78Si9B13 powder particles on the coercive field of composite cores

Fig. 19. Influence of the content of Fe73.5Cu1Nb3Si13.5B9 powder particles on the coercive field of composite cores

Fig. 20. Influence of the content of Fe78Si9B13 powder particles on the saturation induction of composite cores

Fig. 21. Influence of the content of Fe73.5Cu1Nb3Si13.5B9 powder particles on the saturation induction of composite cores

Fig. 22. Influence of the content of Fe78Si9B13 powder particles on the magnetic permeability of composite cores

Fig. 23. Influence of the content of Fe73.5Cu1Nb3Si13.5B9 powder particles on the magnetic permeability of composite cores
Fig. 24. Active power losses and magnetic permeability $\mu_{12000}$ of composites consisting Fe78Si9B13 powder as a function of powder mass share

Fig. 25. Active power losses and magnetic permeability $\mu_{12000}$ of composites consisting Fe73.5Cu1Nb3Si13.5B9 powder as a function of powder mass share

References


