Synthesis, structure and electromagnetic properties of FeCoAl/C nanocomposites

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Received 26 August 2021 ♦ Accepted 10 September 2021 ♦ Published 30 September 2021

Citation: Muratov DG, Kozhitov LV, Yakushko EV, Vasilev AA, Popkova AV, Tarala VA, Korovin EYu (2021) Synthesis, structure and electromagnetic properties of FeCoAl/C nanocomposites. Modern Electronic Materials 7(3): 99–108. https://doi.org/10.3897/j.moem.7.3.77105

Abstract

Magnetic nanoparticles play an important role in rapidly developing advanced branches of science and industry, e.g. fabrication of magnetic storage media, synthesis of ferromagnetic liquids, medicine and chemistry. One problem faced in the usage of magnetic nanoparticles is their high chemical activity leading to oxidation in air and agglomeration. The chemical activity of magnetic nanoparticles stems from the contribution of their large specific surface to volume ratio. Carbon coating of nanoparticles reduces the interaction between nanoparticles. FeCoAl/C metal-carbon nanocomposites have been synthesized using IR pyrolysis of polymer/metal salt precursors. The effect of synthesis temperature (IR heating) in the range from 500 to 700 °C on the structure and composition of the nanomaterials has been studied. We show that the forming particles are the FeCoAl ternary solid solution with a FeCo based bcc lattice. An increase in the synthesis temperature from 500 to 700 °C leads to an increase in the coherent scattering region of three-component nanoparticles from 5 to 19 nm. An increase in the aluminum content from 20 to 30% relative to Fe and Co results in an increase in the size of the nanoparticles to 15 nm but this also entails the formation of a Co based solid solution having an fcc lattice. An increase in the nanocomposite synthesis temperature and a growth of the relative Al content as a result of a more complete carbonization and the structure-building effect of metals reduce the degree of amorphousness of the nanocomposite carbon matrix and lead to the formation of graphite-like phase crystallites having an ordered structure. The effect of synthesis temperature and relative content of metals on the electromagnetic properties (complex permittivity and permeability) of the synthesized nanocomposites has been studied. Synthesis conditions affect the radio absorption properties of the nanocomposites, e.g. reflection loss (RL) in the 3–13 GHz range.

Keywords

FeCoAl nanoparticles, carbon matrix, metal-carbon nanocomposites, IR pyrolysis, X-ray phase analysis, Raman spectroscopy, complex permittivity, complex permeability, reflection loss.
1. Introduction

Magnetic nanoparticles play an important role in rapidly developing advanced branches of science and industry. The use of magnetic nanoparticles shows good promise primarily for the following fields:

- development of magnetic storage media and ferromagnetic liquids [1, 2];
- medicine (transport of medication, contrast agent for magnetic resonance tomography, hyperthermal treatment [3]);
- chemistry (catalysts in petrochemical processes [4]).

The unique physical and chemical properties of iron group metal based alloys originate from their structural properties, electron structure and chemical composition [5–11].

One problem faced in the usage of magnetic nanoparticles is their high chemical activity leading to oxidation in air and agglomeration. The chemical activity of magnetic nanoparticles stems from the contribution of their large specific surface to volume ratio. Carbon coating of nanoparticles reduces the interaction between the nanoparticles but it should be borne in mind that the low degree of crystallinity and graphite shell defects deleteriously affect the protective properties of the coating [12, 13].

Synthesized nanoparticles of iron group metals and alloys without protective coatings are oxidized in air to form oxide films that lead to a significant loss in the ferromagnetic properties. Along with protection, graphite coatings tangibly improve the absorption of electromagnetic waves by metal and alloy nanoparticles. There are two mechanisms underlying this effect: dielectric loss and magnetic loss, with the proper ferromagnetic resonance often occurring at a higher frequency in comparison with that for bulk materials [14].

Another factor allowing one to control the properties of the nanocomposites is the interaction between magnetic particles and the matrix where the magnitude of dipole interaction depends considerably on the textural parameters of the matrix in which the magnetic nanoparticles are dispersed [15].

During the synthesis of nanocomposites the ferromagnetic alloy particles, e.g., FeCo, are protected by shells of a nonmagnetic chemically stable material, i.e., carbon, and are synthesized with different morphologies such as nanocoatings [16–19], nanochips [20], nanotubes [21, 22] and nanofibers [23, 24] with the aim to increase the radio absorption properties of the particles. The main advantage of the use of nanocoated hybrid materials is an increase in the relative complex permeability and dielectric permeability. The mechanisms that increase the complex permeability including the suppression of eddy currents reduce the back reflection and increase the crystallographic magnetic anisotropy which affects the resonance frequency. Different mechanisms such as polarization between the magnetic and dielectric phases and the natural electron relaxation of the dielectric phase also increase the complex permittivity [16–26].

Methods of synthesizing the carbon shells of magnetic nanoparticles include arc discharge [19], hydrothermal treatment [27], magnetron and ion beam sputtering [28], laser pyrolysis of organic compounds [29] and sputtering techniques [30]. The use of carbon as an oxidation protecting shell for magnetic materials has been reported earlier [17–21, 24, 31].

Many of these methods have drawbacks such as incomplete carbon coverage of nanoparticle surface, low graphite formation efficiency and difficult control of carbon coating homogeneity as well as the necessity of an additional heat treatment stage for carbon particle oxidation protection. Therefore an important task is the development of new magnetic material nanoparticle synthesis techniques which can deliver nanoparticles and an oxidation protecting matrix in a single process without the use of a reducing agent.

We developed a method of synthesizing metal/carbon nanocomposites with the use of a system that includes metal salts, a polymer and a solvent by IR heating pyrolysis which is a new and promising trend in the development of new efficient radio absorbing coatings and materials [32, 33]. An important advantage of this method is the possibility of the simultaneous production of multicomponent nanoparticles of FeCoMe (Ni, Sm, Cr, Cu, Al) alloys and a carbon matrix through polymer carbonization. In fact the carbon matrix represents an implementation of the model of a FeCoMe core and a pyrolyzed polymer shell. Cobalt and nickel are quite expensive metals and it is therefore expedient to partially replace them for other metals e.g. Al in multicomponent alloys.

Being aware of FeCoAl/C nanocomposite formation process specific features one can control the magnetic and electromagnetic properties of the materials by varying the elemental composition, structure, morphology and crystalline structure of the multicomponent nanoparticles. Furthermore we gain the possibility of controlling the structure of the carbon matrix which binds the alloy nanoparticles but avoids their agglomeration, oxidation and direct magnetic exchange interaction while retaining a high proportion of the surface.

The importance of this work is dictated by the absence of current literary data on any FeCoAl/C nanocomposite technology.

2. Experimental

FeCoAl/C nanocomposites were synthesized from a precursor consisting of polyacrylonitrile, metal salts and a solvent. Polyacrylonitrile had a molecular weight of 100–250 ths. at. units (powder) produced by oxidation-reduction polymerization. The metal source was iron acetyl acetonate Fe(CH$_2$COCH$_3$)(CH$_3$O)$_2$, 4-aqueous cobalt acetate (Co(CH$_3$COO)$_2$·4H$_2$O) and non-aqueous aluminum chloride (AlCl$_3$).
For achieving a homogeneous initial distribution of the metals in the polymer we made a solution of polyacrylonitrile with metal salts in a common solvent, i.e., dimethylformamide.

After solution drying at 70 °C to a solid residue with a constant weight (hereinafter the precursor) the specimens were subjected to two-stage IR pyrolysis:

- first stage (preliminary air anneal): stepwise heating to 150 °C with 15 min. holding followed by heating to 220 °C with 15 min. tempering (ion air);
- second stage: further vacuum pyrolysis (~200 Pa) at 500 to 700 °C followed by 10 min. tempering, the heating rate being 50 K/min.

X-ray diffraction and phase analyses were conducted at room temperature on a DIFRAY instrument in CrK\textalpha radiation. The experimental results were compared with PDF-4 standards (International Centre for Diffraction Data, ICDD). Based on X-ray diffraction phase analysis data we calculated the average size of the synthesized FeCo alloy nanoparticles using the Debye–Scherrer equations.

Raman spectra were taken on an inVia Raman Microscope spectrometer (Renishaw plc) with 514 nm laser wave excitation.

The complex magnetic and dielectric permeabilities were measured by a resonance method on a rectangular multimode resonator. The HF generator and indicator was a E 8363 B vector network analyzer from Agilent Technologies. The measurements were conducted using instruments and methods developed by the Joint Use Center for Radiological Measurements, Diagnostics and Parametric Research of Natural and Artificial Materials of the Tomsk State University (http://www.ckp.tsu.ru/about/directions/radiophysics).

3. Results and discussion

IR synthesis of polyacrylonitrile and the formation of its carbon matrix involve emission of a significant amount of gaseous products including H\textsubscript{2}, NH\textsubscript{3} and CO which act as reducers for metallic compounds. It should be borne in

![Figure 1](image-url)
mind that reduction occurs in the solid state and therefore metal reduction is an in situ process which may involve atomic hydrogen which forms due to destruction of the main polymer chain during IR heating.

Introduction of a non-magnetic modifying metal (Al) which is often used for the synthesis of various iron based magnetic alloys into the Fe–Co system changes the microstructure and properties of the alloys.

X-ray phase analysis showed that, unlike FeCo/C nanocomponents with an equiatomic content of metals, the formation of a single-phase system does not occur in polycrylonitrile based FeCoAl/C nanocomposites. X-ray patterns of FeCoAl/C nanocomposites synthesized in the 500 to 700 °C range are shown in Fig. 1.

X-ray phase analysis shows that metals or oxides in nanocomposites synthesized at \( T = 500 \, ^\circ\text{C} \) are in an X-ray amorphous state (the average particle size is within 3–5 nm). Two metallic phases having bcc and fcc lattices precipitate at a higher synthesis temperature (600 °C). The good solubility of Al in Fe suggests that bcc FeCo alloy lattice based FeCoAl/C nanocomposites. X-ray patterns of FeCoAl/C nanocomposites synthesized in the 500 to 700 °C range are shown in Fig. 1.

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Increasing the synthesis temperature to 700 °C does not change the phase composition of the nanocomposite. According to the X-ray phase analysis data, bcc lattice FeCoAl nanoparticles co-exist with fcc lattice ones. The X-ray peak of the fcc phase shifts slightly probably due to the dissolution of a small fraction of aluminum in the (Co, Fe) solid solution. The formation of this solid solution is entailed by the presence of iron and cobalt in which aluminum solubilities differ.

It should be noted that the fcc phase peaks are shifted towards lower angles relative to the pure cobalt ones (2θ = 68.02°; 80.48°; 132° (ID: 150806)) suggesting the substitution dissolution of a large quantity of iron and aluminum in cobalt. No aluminum oxides were found; this suggests their X-ray amorphousness or dissolution in other metals with the formation of alloys. The formation of an alloy is further confirmed by the shift of the bcc phase peaks towards lower angles suggesting the dissolution of Al since Al has a larger radius (0.143 nm for Al and 0.126 for iron).

The change in the nanocomposite composition towards an increase in the fraction of aluminum from 20 to 30 rel.% does not lead to any noticeable change in the phase composition of the material but changes the intensity of the peaks of the metallic phases and slightly shifts the peaks relative to the composition Fe : Co : Al = 40 : 40 : 20 (Fig. 2).

Since the radius of the Al atom is greater than that of iron and cobalt the dissolution of aluminum leads to a greater shift of the solid solution fcc phase peaks.

![Figure 2](image-url)  
**Figure 2.** (a) X-ray diffraction patterns of nanocomposites and (b) fragment of X-ray diffraction pattern near 2θ = 65°÷74° for different ratio of metal weights Fe : Co : Al (\( T_{\text{synth}} = 700 \, ^\circ\text{C} \)).

![Figure 3](image-url)  
**Figure 3.** Raman spectra of FeCoAl/C nanocomposites synthesized at different temperatures and metal ratios.
Furthermore the bcc phase peaks shift towards lower angles and this is another indication of aluminum dissolution.

Estimation of the coherent scattering region yielded 10 nm for the fcc phase of the ternary alloy and 15 nm for the bcc phase.

Thus an increase in the fraction of aluminum reduces the average nanoparticle size by 4 nm for the bcc phase and increases the nanoparticle size for the fcc phase.

Heat treatment of polyacrylonitrile triggers a number of chemical reactions, and carbonization produces a carbon material that can be considered as a nanocrystalline one [34]. It was reported that, in the presence of metals, complex carbon structures [35–38] may form which can have a substantial effect on the properties of the nanocomposites obtained.

The Raman spectra of FeCoAl/C metal-carbon composites shown in Fig. 3 demonstrate that with an increase in synthesis temperature the ratio between the D band intensity (I_D) and the G band intensity (I_G) decreases. Furthermore a slight decrease in (I_D/I_G) was observed with an increase in the fraction of Al in the nanocomposites (the Fe : Co : Al ratio was changed from 40 : 40 : 20 to 35 : 35 : 30).

Earlier data [39] demonstrate that a change in the (I_D/I_G) ratio indicates a change in the size of the carbon clusters (L_a) consisting of hexagonal aromatic rings.

The size of these clusters can be estimated using the following equation [40]:

$$I_D/I_G = C(\lambda)/L_a$$  \hspace{1cm} (1)

where C(\lambda) is a coefficient depending on laser wavelength: for laser with \( \lambda = 5.4 \) nm C(\lambda) = 4.4 nm. This estimate showed that L_a with an increase in the synthesis temperature from 500 to 700 °C increases from 25–30 to 40 nm.

In order to study in detail the changes in the microstructure of the carbon matrix as a result of an increase in the synthesis temperature we deconvoluted the Raman spectra into three bands D, G and G_{ac}, where the band G_{ac} peaking at 1520–1540 cm\(^{-1}\) is the contribution of amorphous carbon clusters.

It can be seen from Fig. 4 that the greatest structural transformations in the test materials occur at 500 to 600 °C. In accordance with earlier data [41] the trend of decreasing the (I_D/I_{Gac}) ratio indicates a decrease in the size of amorphous carbon clusters.

Each band was described using the Lorentz distribution.

The results showed that with an increase in the synthesis temperature of FeCoAl/C nanocomposites the sizes of the crystalline clusters in the microstructure of the carbon matrix decrease, i.e., the amorphousness degree of the material decreases.

The structural transformations in the nanocomposite matrix and the changes in the size and composition of nanoparticles with an increase in the synthesis temperature entail tangible changes in the electromagnetic properties of the materials.

Data on the effect of synthesis temperature on the dielectric and magnetic properties of the nanocomposites in HF fields are shown in Fig. 5.

The data shown in Fig. 5 b suggest that the \( \mu^* \) component of the complex permeability increases significantly with an increase in the synthesis temperature. Taking into account that this component describes the fraction of absorbed radiation, an increase in this parameter from 0.04 to 0.12 leads to an increase in the magnetic loss. The curve has a peak at ~11 GHz which seems to originate from an increase in FeCoAl alloy nanoparticle sizes and hence an increase in the magnetization of the nanocomposite. The real part of the complex magnetic permeability also increases by approx. 10% with an increase in the
nanocomposite synthesis temperature. Coupled with the changes in the imaginary part of the complex magnetic permeability this causes an increase in the magnetic loss tangent in the test frequency range.

The dielectric permeability and the magnetic dielectric loss tangent (Fig. 5a and c) also increase with the synthesis temperature. The growth is the greatest in the 500 to 600 °C range which is in agreement with the X-ray phase analysis data and the conclusions from the Raman study.

From the practical application viewpoint the fabrication of radio absorbing components of structures on the basis of the nanocomposites synthesized in this work having the smallest weight and dimensions is the most efficient for the frequencies of natural ferromagnetic resonance since the magnetic loss tangent is the largest in this frequency range.

Calculations showed that the optimum layer thickness for the synthesis of absorbing components on the basis of FeCoAl/C nanocomposites differs substantially for specimens synthesized at different temperatures (Fig. 6). For example, for nanocomposites synthesized at 700 °C the optimum thicknesses are 1.5 and 4.2 mm. The reflection loss coefficient (RL) is as high as –36 dB (absorption coefficient 0.984). For nanocomposites synthesized at 500 and 600 °C the optimum layer thicknesses are at higher frequencies compared with the range considered (3–13 GHz).

The calculation results suggest (Fig. 6) that the specimens synthesized at 500 and 600 °C have low efficiencies over the entire range of frequencies and thicknesses since their absorption coefficient is within 57% (RL ≥ 7.2 dB). For FeCoAl/C nanocomposite synthesized at 700 °C having the optimum thickness the maximum absorption is

Figure 5. Frequency dependences of (a) complex permittivity and (b) permeability and tangents of (c) dielectric and (d) magnetic loss for different FeCoAl/C nanocomposite synthesis temperatures.

Figure 6. Optimization of absorbing layer thickness.
Figure 7. Frequency dependences of reflection coefficient of FeCoAl/C nanocomposites synthesized at different temperatures $T_{\text{synth}}, ^\circ \text{C}$: (a) 500, (b) 600 and (c) 700.

achieved at 13.8 GHz for an absorbing layer thickness of 1.5 mm and with an increase in the nanocomposite layer thickness the minimum reflection (maximum absorption) shifts towards lower frequencies and reaches –17 dB at a 7 mm thickness.

Calculated data for reflection loss taking into account nanocomposite position on metal are shown in Fig. 7. We also studied the effect of metal ratio in the precursor on the electromagnetic and radio absorption properties of the FeCoAl/C nanocomposites. The studies showed that the dielectric loss tangent and the dielectric permeability components decrease with an increase in the fraction of aluminum (Fig. 8 a and c) which seems to be caused by an increase in the overall electrical conductivity of the nanocomposite.

The magnetic loss is almost constant although the formation of larger cobalt based solid solution particles makes the spectrum more uniform in the 3–13 GHz frequency range, growing from 0.03 to 0.1 rel.u.

Calculated data on the reflection loss taking into account nanocomposite position on the metal are shown in Fig. 9.

The minimum reflection (RL = –14.3 dB) and the maximum absorption (80.7%) can be achieved at an absorbing layer thickness of 3 mm which is inferior to other nanocomposites and to FeCoAl/C nanocomposite with the 40 : 40 : 20 metal content ratio.

Thus the most promising FeCoAl/C nanocomposites are those synthesized at $T_{\text{synth}} = 700 ^\circ \text{C}$ and having the metal content ratio Fe : Co : Al = 40 : 40 : 20.

4. Conclusion

The possibility of synthesizing nanocomposites containing nanoparticles of the FeCoAl ternary alloy (solid solution) using IR pyrolysis of metallic precursors was demonstrated. Nanoparticles of fcc Co and bcc FeCo based solid solutions form with an increase in the synthesis temperature leading to an increase in the size of the coherent scattering region for alloy nanoparticles from 5 to 19 nm due to agglomeration and coalescence of the metals in the course of reduction. An increase in the fraction of aluminum entails a decrease in the size of FeCoAl nanoparticles having a bcc lattice (15 nm) and an increase in the fraction of nanoparticles having an fcc lattice. An increase in the synthesis temperature and the fraction of aluminum in the nanocomposite lead to the formation of a more ordered nanocomposite carbon matrix which shows itself in a higher degree of crystallinity and a decrease in the amor-
Matrix structurization leads to an increase in the dielectric loss due to the formation of crystalline inhomogeneities and a decrease in the amorphousness degree. In the meantime an increase in the size of nanoparticles entails an increase in the magnetic loss upon interaction with microwave radiation in the 3–13 GHz range. Furthermore the interaction between the nanocomposite matrix and alloy nanoparticles increases the permittivity thus providing for a deeper penetration of electromagnetic waves into the material and more efficient energy dissipation. The most promising radio absorbing materials are FeCoAl/C nanocomposites synthesized at 700 °C. Their reflection loss is from −17 to −31 dB which corresponds to a 86 to 97.2% absorption index.

Figure 8. Frequency dependences of (a) complex permittivity and (b) permeability and tangents of (c) dielectric and (d) magnetic loss for different metal ratios in precursor.

Figure 9. Frequency dependences of reflection coefficient of nanocomposites for different metal content ratios in precursor Fe:Co:Al: (a) 40:40:20; (b) 35:35:30.
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