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MICROSTRUCTURE OF MAGNETITE DOPED ELASTOMERS INVESTIGATED BY SAXS AND SANS

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Балашою М. и др. Исследования микроструктуры магнитных эластомеров на магнетите методами МУРР и МУРН

Мы представляем последние результаты исследования магнитных эластомеров методом малоуглового рассеяния рентгеновского и нейтронного излучения. Образцы были получены в процессе полимеризации диметилсилоксилана с магнитной жидкостью с частицами Fe₃O₄ в магнитном поле и в отсутствие поля и были исследованы методами рентгеновской дифракции, МУРН и МУРР. Пик, присутствующий в области больших переданных импульсов Q кривых МУРН, полученных с магнитных эластомеров, был обработан с использованием для фитирования Pseudo-Voigt функции. Было найдено, что частицы Fe₃O₄ приводят к изменению локальной структуры эластомера, т. е. уменьшению квазикристаллической фазы и микронапряжений, индуцированных в матрице. Средний размер частиц, полученный из измерений МУРН, согласуется со средним размером частицы, определенным с помощью рентгеновской дифракции и МУРР. Разница в интенсивности малоуглового рассеяния нейтронов вследствие дополнительного рассеяния на образце, полимеризованном в нулевом магнитном поле, показывает существование магнитных корреляций в области 0,005 < Q < 0,02 Å⁻¹.

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Balasoiu M. et al. Microstructure of Magnetite Doped Elastomers Investigated by SAXS and SANS

We present some recent results on the SAXS and SANS investigations of the magnetic elastomers. Samples were obtained by polymerizing dimethylsiloxane with Fe₃O₄ ferrofluid in a magnetic field and without magnetic field and characterized by XRD, SANS and SAXS. The SAXS maximum at large angles, obtained from magnetic elastomers, was fitted by using a Pseudo-Voigt function. It was found that the Fe₃O₄ particles lead to a change of the local structure of elastomer, meaning the decrease of the quasi-crystalline phase and of the microstrains induced in the matrix. The average particle size obtained from SANS measurements agrees well with the average particle size determined by XRD and SAXS analysis. The splitting of the SANS intensity owing to an additional scattering in zero field reflects the existence of magnetic correlations inside the *Q* region of 0.005 < Q < 0.02 Å⁻¹.

The investigation has been performed at the Frank Laboratory of Neutron Physics, JINR.

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INTRODUCTION

Magnetic elastomers belong to a specific class of so-called smart materials because they can respond to changes in their environment. They are composed of magnetic particles and a low-permeability matrix. Applying an external magnetic field, a structure will be formed inside the material or the structure embedded in the material will be changed. Combination of magnetic and elastic properties leads to different phenomena which are exhibited in a variable magnetic field [1–3]. It opens new possibilities for technological applications as: 1) magnetoelastic composite with particles made of magnetostrictive hard or soft ferromagnetic material [4, 5]; 2) magnetorheologic elastomers for application in airplane and car industries as actuators or antifriction components [6]; 3) heat-shrinkable elastic ferromagnets with variable magnetic and conductive properties [7].

Other applications that utilize magnetic polymer nanocomposites are currently emerging at a high rate. Examples include magnetic actuation in microelectromechanical systems (MEMS) and medical devices, thermal actuation through electromagnetic power harvesting, and magnetically actuated morphing structures [8].

Such composites are quite new and understanding of the behavior of these materials depending on the composition, external conditions, and the synthesis processes is intensively under investigation [9].

The magnetoelastic properties of composites are not the bare sum of the elasticity of the polymer and the stiffness and magnetic properties of the filler, but are the result of a complex synergy of several effects, referring to different length scales and detectable by different techniques.

Many studies of the observed reinforcing effect from magnetic fillers have approached the problem from a magnetomechanical point of view and investigated the microscopic properties through the study of the magnetoelastic responses of the composite [1-3, 10]. Less well-understood, however, is the effect of the polymer-filled interaction with the local distribution of polymer around filler on a submicroscopic length scale.

Work making use of scattering techniques as neutron and X-rays is rarer but if used, is based on the scattering vector dependence or exploits the full advantage relating the intensity to the volume of the scattering particle [11, 12].

1. EXPERIMENTAL

The studied samples, obtained at the Department of Electricity and Magnetism, West University of Timisoara [13–15], are composed of oil-based 7.7% particle volume concentration Fe₃O₄ ferrofluid with oleic acid as surfactant, embedded in a polymer matrix formed from dimethylsiloxane, dibutyltindilaurate benzyl silicate, polymerized in zero field (sample A) or in an applied magnetic field of H = 108.28 kA/m (sample B) [16].

In order to determine the phase composition, lattice microdistortions and average size of coherent length (average size of mosaic blocks), the samples were investigated by XRD. The XRD measurements were performed using DRON diffractometer, $CoK\alpha$ radiation. The XRD data were handled using Ceck Cell and Rietveld software.

The samples were studied using small-angle X-ray (SAXS) and neutron (SANS) scattering methods. The SAXS experiments were performed at Rigaku spectrometer, using a pinhole camera (Molecular Metrology SAXS System) attached to a microfocused X-ray beam generator (Osmic MicroMax 002). The camera was equipped with a multiwire, gas-filled area detector with an active area diameter of 20 cm (see a schematic representation of small-angle diffraction setup in Fig. 1). Two experimental setups were used to cover the Q range of 0.007-1.1 Å⁻¹ (Fig. 3).



Fig. 1. Representation scheme of the small-angle diffraction setup

The SANS experiments were carried out on the YUMO diffractometer [17] at the IBR-2 pulsed reactor, JINR, Dubna and on SANS-1 [18] installed at Geesthacht, Germany in the Q range of about 0.003-0.3 Å⁻¹.

2. SMALL-ANGLE SCATTERING ELEMENTS

In the small-angle neutron scattering experiment, $d\Sigma/dQ\Omega$, the differential scattering cross section is measured as a function of the momentum transfer, $Q = (4\pi/\lambda)\sin(\theta/2)$. λ is the neutron's wavelength and θ — the scattering angle.

Typically, scattering experiments at a given value of Q probe a length scale D, where D is the dimension of the inhomogeneities producing the scattering:

$$D = 2\pi/Q.$$
 (1)

The SANS range is usually defined by $0 < Q < \pi/D$. For dilute systems of randomly oriented particles (neglecting size polydispersity) in the region of small Q values ($QR_g < 1.2$), Guinier's law is applied and the scattering is given by the relation:

$$d\Sigma(Q)/d\Omega = d\Sigma(0)/d\Omega \exp\left(-R_a^2 Q^2/3\right),\tag{2}$$

where, $d\Sigma(0)/d\Omega$ is the scattering at zero angle and R_g is the radius of gyration of the particle, i.e., the root-mean-square distance of all scattering elements from the centre of gravity. If this approximation holds, the plot $\ln I(Q)$ versus Q^2 should show a linear dependence in the region of small Q. The characteristic size of a particulate system can be easily obtained within this approximation. For instance, for spherical particles, one can relate the radius of particle R to the radius of gyration by $R = (5/3)^{1/2} R_g$ [19].

3. DISCUSSIONS

From XRD data results that the elastomer was doped with Fe_3O_4 particles about 10 nm average size, the elastomers contain practically the same concentration of Fe_3O_4 , but with different average size of coherent blocks and microstrains.



Fig. 2. Observed and calculated diffractogram (upper-side); difference between observed and calculated diffractogram (bottom-side) for Fe_3O_4 particles

The data were handled with PowderCell, to obtain the lattice constants, average size of crystalline blocks and microstrains. It was established that both samples contain a small amount of foreign phase (Fig. 2).

To obtain exact lattice constant, the average size of mosaic blocks and microstrains of magnetite, we used a Rietveld program. In agreement with the literature data, magnetite crystallizes in GS 227, with a = 8.38 Å; atomic position in unit cell is given in Table 1.

Table 1. Atomic position (x, y, z), occupation and temperature factor (B)

Ion	Atomic coordinates			Occupation	В
	x	y	z	Occupation	D
Fe ³⁺	0.125	0.125	0.125	1.0	0.25
Fe ³⁺	0.500	0.500	0.500	0.5	0.35
Fe ²⁺	0.500	0.500	0.500	0.5	0.35
0^{2-}	1.000	1.000	0.754	1.0	1.32

Data concerning SAXS measurements indicated a strong difference of the structure of doped and undoped with Fe_3O_4 particles samples. We had observed the systematic appearance of a Bragg diffraction peak in SAXS plots in the region of large Q values (near Q = 0.9 Å⁻¹; see Fig. 3). The lattice parameter of quasicrystal was obtained by fitting the observed maximum with a Pseudo-Voigt profile function (Fig. 4).



Fig. 3. SAXS experimental curves from samples A, B and simple elastomer obtained at Rigaku spectrometer in function at IMC, Prague



Fig. 4. Fit of the XRD data using a Pseudo-Voigt profile function

We suppose that the elastomer is fragmented into small particles. Doping with Fe_3O_4 particles leads to a significant change of the local structure of elastomer, meaning the decrease of the of quasi-crystalline phase concentration (Fig. 3) and average size of the crystalline blocks (Table 2). This maximum was fitted using a Pseudo-Voigt profile function (Fig. 4).

The average size of elastomer blocks and the ordering distance decrease when magnetic field is applied during the polymerization process (Table 2).

Figure 5 shows the curves of the neutron scattering cross section for the samples of the Fe₃O₄ elastomer polymerized in zero magnetic field (sample A) and in a magnetic field of H = 108.28 Am⁻¹ (sample B).

Following the data evaluation of [9] we use the fact that in the case of sample B polymerized in magnetic field, the magnetization of most of the particles is aligned in the direction of the applied field. In this case, the forced common magnetic alignment extends to such large distances that it cannot be resolved any more by SANS.

Table 2. Variation of gravity center position (θ), average size of crystalline blocks (Δ) and ordering distance (D) in the elastomer matrix

Sample	θ	Δ , nm	D, nm
Elastomer matrix	11.56 ± 0.02	2.73 ± 0.04	0.77 ± 0.001
Elastomer with Fe ₃ O ₄ particles	11.71 ± 0.01	2.38 ± 0.13	0.77 ± 0.001
(polymerization without magnetic field)			
Elastomer with Fe ₃ O ₄ particles	11.41 ± 0.05	2.73 ± 0.14	0.77 ± 0.01
(polymerization with magnetic field)			



Fig. 5. SANS scattering from elastomer A and B; SANS scattering from elastomer A (scattering from pure elastomer subtracted); curves obtained at SANS-1 in function at GeNF

The scattering then originates from nanometer sized particles surrounded by particle boundaries of different (nuclear and magnetic) density.

The average particle size obtained from these measurements agrees well with the average particle size determined by XRD and SAXS analysis.

Table 3. Radius of gyration and particle radius $(R = (5/3)^{1/2}R_g)$ obtained from SAXS and SANS measurements, respectively

Sample	R_g /SAXS	R_g /SANS	R, nm
Elastomer with Fe ₃ O ₄ particles	1.9 ± 0.01	2.29 ± 0.05	2.45 ± 0.01
(polymerization without magnetic field (A))			2.95 ± 0.06
Elastomer with Fe ₃ O ₄ particles	2 ± 0.01	2.3 ± 0.05	2.58 ± 0.01
(polymerization with magnetic field (B))			2.96 ± 0.06

The corresponding scattering curves of the neutron scattering cross section for the samples polymerized in zero magnetic field and in a magnetic field lie closely together and are parallel for large Q values; for smaller Q values a

splitting is observed. The splitting of the SANS intensity owing to an additional scattering in zero field reflects the magnetic correlations inside the Q region of 0.005 < Q < 0.02 Å⁻¹ existence.

Such correlations were also found in other SANS studies [20-26].

CONCLUSIONS

It was obtained that: the profile of diffraction maximum belonging to magnetite can be fitted by using a Pseudo-Voigt function. We found that magnetite particles are nanometer size (9.8 nm). The samples contain small amount of foreign phases. We supposed that the elastomer is fragmented into small particles ordered at short distance. Doping with Fe_3O_4 particles leads to a significant change of the local structure of elastomer, meaning the decrease of the microstrains in the matrix.

The average particle size obtained from SANS measurements agrees well with the average particle size determined by XRD and SAXS analysis.

The corresponding scattering curves of the neutron scattering cross section for the samples polymerized in zero magnetic field and in a magnetic field lie closely together and are parallel for large Q values; for smaller Q values a splitting is observed. The splitting of the SANS intensity owing to an additional scattering in zero field reflects the existence of magnetic correlations inside the Q region of 0.005 < Q < 0.02 Å⁻¹. This shows that the magnetic correlations are not confined to single particles, but extend across the interfaces and result in a common, localized magnetic alignment of many grains.

In future we intend to perform systematic investigations concerning structure, magnetic and elastic properties of elastomers doped with magnetic nanoparticles.

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