The common practice of using iron nanoparticles in human and veterinary medicine as well as their potential use microelement-based medicines determine the need for studying the impact that nanoparticles have on the exchange of chemical elements in the body. The study involved a Wistar rats model using iron nanoparticles (nanoFe), obtained through high-temperature condensation \((d = 80 \pm 5 \text{ nm})\). The study on genetically engineered luminescent strain E. coli K12 TG1 had a pre-installed non-toxic concentration of nanoFe. Atomic emission and mass spectrometry showed the presence of 25 chemical elements in the animals’ liver after seven nanoFe intramuscular injections had been given to them. The experiment revealed no disturbance in the liver microstructure. However, an investigation into the dynamics of transaminases (alanine transaminase (ALT), aspartate transaminase (AST)) revealed an increase in their activity. On Day 1 of the experiment the LDH activity went 116.3\% up \((p < 0.001)\) to go down gradually within 21 days. Intramuscular nanoFe injections came along with certain alteration in the exchange of chemical elements. A single dose of iron nanoparticles caused, in the first seven days, depletion of the liver and its saturation with toxic elements. On the first day after the first injection this was manifested through an increase in the concentration of Pb by 20.0\% \((p < 0.05)\), Sn by 33.3\% \((p < 0.05)\), Sr by 66.67\% \((p < 0.01)\). The most significant adaptive changes in the toxic elements exchange of were observed for Al and Sr. The iron content in the liver decreased on Day 7 after the first injection by 19.35\% \((p < 0.05)\), Day 2 by 28.9\% \((p < 0.05)\), Day 3 by 7.01\%, Day 7 by 16.79\% \((p < 0.05)\) compared to the controls. The pool of the macronutrients Ca, K, Mg, Na, P (the sum of the substance amount, mole) was found to vary through the experiment by 4.1–10.4\%. Reduction of calcium concentration one day following the first injection (in comparison to the controls) was 6.81\%; on Day 7 after the second injection – by 18.58\% \((p < 0.05)\); after the third and the seventh injections – by 6.1% and 12.4\% \((p < 0.01)\), respectively. Various studies suggest that there is a need for additional correction of the elemental composition in diets against iron nanoparticles injections.

Keywords: Element Status, Iron Nanoparticles, Rats
The currently available preparations based on this microelement\textsuperscript{11}, etc. Even though iron nanoparticles and its compounds are practically used more and more often there is still no proper understanding of the subcellular impact that these structures have. There facts reporting certain side effect of nanoparticles, which are manifested as disturbed immunity\textsuperscript{12, 13}. Iron and its compounds injections, local or systemic, trigger the development of oxidative stress, which leads to an acute inflammation response\textsuperscript{14} and are accompanied with toxicosis in animals\textsuperscript{15}. These facts imply comprehensive study of the biological effects caused by the agents based on such nanoparticles. This appears especially urgent in view of the fact that the nanoparticles are promising as commercial microelement agents and possess a number of advantages if matched against mineral salts and organic forms. Selenium nanoparticles, in particular, are less toxic and more effective if compared to selenite and other preparations\textsuperscript{16-19}.

This explains the interest taken in the impact that iron-containing nanoparticles introduced into the body have on the exchange of other chemical elements. The purpose of this present study is to investigate the effect of iron nanoparticles on microelement exchange in rats.

**MATERIALS AND METHODS**

**Obtaining and notification of iron nanoparticles**

Iron nanoparticles (nanoFe) were obtained through high-temperature condensation on a Migen-3 machine\textsuperscript{20}. The nanoparticles were spherical in shape, sized 80±5 nm, Z-potential – 15±0.2 mV. The material notification of the preparations included scanning and transmission electron microscopy using the machines like JSM 7401F, JEM-2000FX (JEOL, Japan); X-ray phase analysis on the diffractometer DRON-7. The AFM investigation was done on the microscope SMM-2000 (JSC PROTON-MIET, Russia). Through the scanning there were used the cantilevers MSCT-AUNM (Park Scientific Instruments, NOA) with a beam stiffness of 0.01 N/m and a needle curvature radius of 15-20 nm. The quantitative morphometric analysis of the obtained images was performed with the actual software for the microscope.

All the experiments were done in triplicate and processed by variation statistics using the software package Statistika V10 RUS). The biological activity and the nanoFe toxicity thresholds were detected through bioluminescence inhibition method. NanoFe samples were prepared at a concentration of 4 mole/l on physiological solution and were ultrasound-treated for 30 minutes (ultrasonic disperser UZDN-2T, (Russia) att-35 kHz, N 300 W, and A-10 ia). To evaluate the effect of various nanoFe dosages, the resulting suspensions were used to prepare ten serial double dilutions. The genetically engineered luminescent strain E. coli K12 TG1 was used; this strain was engineered to constitutively express the luxCDABE genes of the natural marine microorganism Photobacterium leiognathi 54D10 and was produced by Immunotech (Moscow, Russia). In prior studies, the strain Echerichia coli K12 TG1 was restored by the addition of chilled distilled water. The suspension of bacteria was maintained at +2-4ºÑ for 30 min, after which the temperature of the bacterial suspension was brought to 15-25ºC.

The inhibition of bacterial luminescence was tested by placing the cells in 96-well plates containing the test substance and the suspension of luminescent bacteria in a 1:1 ratio. Subsequently, the tray was placed in the measuring unit of an Infinite PROF200 microplate analyzer (TECAN, Austria), which dynamically registered the luminescence intensity for 180 min at intervals of 5 min.

The effects of the nanomaterials on the intensity of bacterial bioluminescence (I) were evaluated using the formula:

\[ I = \frac{I_{\text{km} \text{in}} \times I_{\text{o} \text{min}}}{I_{\text{km} \text{in}} \times I_{\text{o} \text{min}}} \]

where \( I_{\text{km}} \) and \( I_{\text{o}} \) are the illumination intensities of the control and experimental samples, respectively, from the 0-th and \( n \)-th minutes of measurement. Three threshold levels of toxicity are taken into account:

1. less than 20 – sample is “non-toxic” (luminescence quenching ≤ 20 %);  
2. from 20 to 50 – sample is relatively toxic (luminescence quenching 50 %);  
3. equal to or greater than 50 – sample toxic (luminescence quenching ≥ 50 %).

**In vivo methods**
The research in vivo was conducted on male Wistar rats, 150-180 g. The animals were divided into two groups (pair-analogue method used) (n=50). The animals were kept on natural and well-balanced diets typical of rodents. The animals once a week were injected with iron nanoparticle in femoral group of muscles in dosage of 2.0 mg/kg of weight (for 7 weeks, with a total of 7 injections). The control group animals were injected with sterile physiological solution (200 mcl/head). The injection sites were chosen at distances, and in view of the muscle regeneration terms and respective recommendations; repeated injection in the same area was given no earlier than 3 weeks after. The experimental research on animals was done following the instructions set by the respective Russian Regulations (1987) and The Guide for the Care and Use of Laboratory Animals (National Academy Press Washington, D.C., 1996).

The nanoFe injection was prepared through mixing nanoparticles with physiological solution (200 mcl) after which the preparation was sterilized with UV to be further treated with ultrasound for 30 minutes (ultrasonic disperser UZDN-2T, (Russia) atf-35 kHz, N 300 W, andA-10 ia). The animal biosubstrates were taken at slaughter, which was performed (n=3) by decapitation under Nembutal narcosis 1 and 7 days after each injection.

The element composition of the biosubstrates was studied with atomic emission and mass spectroscopy at the experimental laboratory of the Center for Biotoc Medicine, Moscow, Russia (Registration Certificate of ISO 9001: 2000, Number 4017 – 5.04.06). The biosubstrate ashing was performed with the microwave decomposition system MD-2000 (USA). The element content was determined with the mass-spectrometer Elan 9000 and the atomic emission spectrometer Optima 2000 V (PerkinElmer, USA).

For light microscopy pieces of liver were fixed in a 10 % formalin solution. The paraffin sections (5-6 mcm) were stained with Meyer’s hematoxyline-eosin. Iron was detected in the studied organ through Pearls reaction.

RESULTS

In vitro study results

The results obtained allowed describing the dynamics of bacterial bioluminescence inhibition through time, as well as demonstrating the link between the registered effects and different concentrations of nanoFe.

NanoFe preparation in a dosage of 0.5 mole/l (28 g/l) resulted in 50 % bacteria luminescence quenching 60 minutes after the contact, if compared with the controls, with complete suppression of bioluminescence 160 minutes after the contact. When taken in concentrations like 0.25 mole/l (14 g/l) and 0.1 mole/l (5.6 g/l) nanoFe revealed a weak toxic effect resulting in a 30 % bioluminescence quenching 80 minutes after the contact. NanoFe concentration lying within the range of 0.05-0.000781 mole/l (2.8-0.044 g/l) showed lack of substantial impact on the microorganism bioluminescence. The data was used for detecting non-toxic nanoFe dosages in intramuscular injections.

In vivo study results

A study of the mineral composition in the animals’ tissues conducted within 49 days showed certain changes in the chemical element exchange in rats, which was due to nanoFe impact. A single injection of nanoFe within the first seven days revealed that the liver had a reduced level of Mg (by 28.38%; p<0.05), K (by 22.45%; p<0.05). The concentration of the toxic elements (tin monoxide, lad, and strontium) went up. The first nanoFe injection was associated with a reduced iron concentration in the liver (by 18.25 %) (Figure 1 A, B). The Pearls qualitative histochemical reaction in the liver produced positive result only on Day 14 after 1 injection.

The experiment revealed no disturbance in the liver microstructure. However, an investigation into the dynamics of transaminase activity (AST, ALT) showed an increase in their activity. ALT activity on Day 3 after the first injection went above the controls by 69.5 % (p<0.05) (Figure 2). On Day 7 the ALT activity in the experimental group went down exceeding the control values by 29.8%. On Day 14 and Day 21 after the first injection the blood transaminase
Fig. 1. Difference in the mineral element concentration in the liver of the animals, control group, 1 day after (A), and 7 days after (B) the first nanoFe injection; dosage – 2 mcg / kg, %

Fig. 2. ALT Levels in blood serum in rats after the 1st injection of nanoFe, dosage – 2 mg/kg, U/l
Fig. 5. Difference in the mineral element concentration in the liver of the animals, the experimental group compared to the control group, Day 7 after the 7th nanoFe injection; dosage – 2 mcg / kg, %

The blood transaminase activity in the rats assessed by AST went up on Day 1 of the experiment by 27.9% (Fig. 3). Later on it went down compared to the control values – by 24.4% on Day 3 and by 29.6% - on Day 7. Through the second and the third weeks of the experiment, the AST activity was above the controls – Day 14 – by 123% (Ð < 0.01), Day 21 – by 115%. The AST activity growth describes the nanoparticle effect as cytotoxic.

The adaptive changes in the chemical activity in the experimental group was below that in the control group by 49.3% and by 30.7% (Ð < 0.05).

Fig. 3. Levels of AST in blood serum in rats at injecting nanoFe, dosage 2 mg/kg, U/l data presented as: mean (X) ± standard error of the mean (SE), * - results are statistically significant (p<0.05), ** – results are statistically significant (p<0.01)

Fig. 4. Liver. Negative Pearls reaction on Day 7 after the 7th injection. Magnification – 600
element exchange in the experimental group were to be observed through the entire trial. The most prominent changes were seen in iron exchange. The level of iron in the liver went down on Day 7 after the 1st injection by 19.35 % (p<0.05), after the 2nd injection – by 28.9 % (p<0.05), after the 3rd – by 7.01 %, after the 7th – 16.79 % (p<0.05) compared to the controls.

Based on the presence of a focal proliferation after the 5th injection, we can talk about activated performance in the Kupffer cells inside some liver acini. However, by the end of the study the Kupffer reaction was negative (Figure 4).

The study showed an increase in the iron content in the blood serum in the animals involved - by 20.3% (p<0.001) after 1 day; by 23.5% (p<0.001) – 7 days, and by 18.8% (p<0.001) 21 days after the injection.

Significant changes were observed in the exchange of essential microelements Cr, Cu, B, Zn already on Day 1 after the first injection (Figure. 1). In particular, the levels of Zn on Day 1 after the injection went down by 16.57%. The next three days revealed a steady increase in the Zn content. The dynamics in the Zn concentration may be linked to an increased level of lactatedehydrogenase (LDH), which is a zinc-containing enzyme. On Day 1 of the experiment the LDH activity went up by 116.3 % (p< 0.001) compared to the controls. Further on, on Day 7 and Day 21 the values in the experiment and the controls got equal remaining high at 693 IU/l on Day 7, and went down back to the initial values of 377.3 IU/l in the controls and 351.5 – in the experimental group on Day 21.

Intramuscular injections of nanoFe were associated with changes in the exchange of toxic elements in the liver. On Day 1 after the 1st injection it manifested itself through an increase in the concentration of nearly all the elements under study: Pb by 20.0 % (p<0.05), Sn by 33.3% (p<0.05), Sr by 66.67% (p<0.01). The adaptive changes in the toxic element exchange revealed themselves in a relative decrease in their levels in the animals’ liver within the first three weeks after the injection.

The most significant decrease was observed in Al – 60.5% and Sr – 76.7% compared to the control values. Seven consecutive injections of nanoFe resulted in different changes in the toxic elements content. On Day 7 after the 7th injection the level of Al went up beyond that in the control group by 17.39% (p<0.05), Cd – by 40.00% (p<0.001), Pb – by 25.0% (p<0.001). At the same time the concentration of tin and strontium, on the contrary, went down against the control values by 60.0 % (p<0.0001) and 26.67% (p<0.0001), respectively.

The content of arsenic went up significantly after the 2nd injection, while the concentration difference with the third injection was by 210.3 % (p<0.05). The Ni concentration at single injection of nanoparticles went down by 33.3 % (p<0.01) compared to the control group; in case of multiple injections, however, there was an opposite effect – an increase by 14.29 %. The Cu concentration in the experimental group went down after the 1st injection by 17.8 %, after the 2nd injection – by 8.5 %, after the 3rd injection – by 4.2 %, and a 7 % (p<0.05) increase was observed following the 7th injection.

**DISCUSSION**

The common practice of using iron nanoparticles as well as their potential use as microelement preparations explain the need for investigating the impact caused by the nanoparticles on other chemical elements exchange. This need is due to the synergy and the antagonism of these elements. The study presented here serves evidence to this fact. Intramuscular injections of nanoFe were associated with changes in the chemical element exchange in the animals. Looking into the reasons behind these changes

While looking at the reasons behind these changes it must be noted that our study does not offer a description as to a prominent toxic
effect of nanoFe. This is confirmed through our study involving the model E. coli K12 TG1, at assessing the morphology of the liver, and in earlier works on multiple injections of nanoparticles23.

The choice of liver as the model is due to the fact that it is one of the major depots for microelements, which are able of being involved into exchange with plasma for several hours24.

In our studies multiple injections of nanoFe came along with reduced content of this element in the liver, which was rather natural. Iron concentrations in biological fluids are known to be subject to tough regulation25. Excess of iron may result in generation of active forms of oxygen26, 27.

Iron injected intramuscularly is deposited at the injection site to be further released gradually from the depot causing increases in hemoglobin, serum iron, and ferritin28.

The development of homeostatic reaction to nanoFe injection was associated with a gradual decrease of selenium content in the liver. In particular, a week after the first injection the content of selenium in the animals’ liver went beyond the control levels by 7.25%. Later on it went down steadily, compare to the controls, on Day 7 after the 2nd injection (by 1.16%) and the 3rd injection (by 6.1% (p<0.05)), after the 7th injection (by 17.57% (p<0.01)). Selenium consumption was detected through its involvement into compounds of selenium-proteins needed for protecting cells from oxidative materials29, 30, 31 as well as from heavy metals against reducing iron concentrations, which is compatible with the earlier obtained data on antagonism of these elements32. The reasons behind this include competition for common transport proteins for iron and other bivalent metals in the intestines33, 34.

Earlier it has been shown that additional introduction of iron into blood is associated with ferroportin synthesis35. Obviously, the capacity of ferroportin to transport other metals, too, including Cd 36, 37 might lead to certain change in their total pool in the body.

Oxidative stress induced by nanoFe damages cells38. This is why there were earlier reports about pathological processes taking place in the liver in case of introduction of nanoparticles of Fe, Np, Zn39.

Our research shows that the liver reveals no pathomorphological processes evaluated through light visualization However, when evaluating transaminase dynamics (AST, ALT) there was an increase in their activity detected, which might be a consequence to damaged cell membrane.

Similar results were obtained when injection rats with iron oxide nanoparticles (dosage 10 mg/kg). The nanoparticles injections were associated with increased levels of aspartate transaminase (AST), alanine transaminase (ALT), alkaline phosphatase (ALP), and gamma-glutamyl transferase (γGT). However, histological analysis of the liver, kidneys, spleen, lungs, brain, and heart revealed no typical damage40.

CONCLUSION

Intramuscular injections with iron nanoparticles are associated with significant changes in the animals’ element status already on the first day after the injection and remain there for up to three weeks. Along with vital elements, the effect of iron nanoparticles expands to cover toxic elements as well. Studies suggest that it is advisable to perform additional correction of the element composition in diets in case of iron nanoparticles injections.

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