Comparative effects of intramyocardial autotransplantation of different bone marrow cells upon outcomes of experimental myocardial infarction in rabbits

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Summary

Over the last decade, different cell therapy options have been tested to improve treatment of myocardial infarction (MI), including cardiomyoplastics. Autologous bone marrow is a widely used source of therapeutic cell preparations, i.e., freshly isolated mononuclear cell fraction (MF), or cultured multipotent mesenchymal stromal cells (MMSC). The aim of our pilot study was to compare short- and long-term consequences of intramyocardial MF and MMSC transplantation upon clinical course and outcome of experimental MI. Materials and methods: The experiments were performed with male Chinchilla rabbits of 2.8±0.2 kg weight. MI was modelled by ligation of anterior descendent left coronary artery. Mononuclear cell fraction was obtained from the bone marrow aspirate. MMSC cell culture was grown by MF cell passages in MEm medium with 10% fetal calf serum. Intramyocardial injections of MF or MMSC suspensions were performed into 6 points of the infarcted area. The surviving animals were divided in four groups, each consisting of 13 rabbits: group 1 (controls), group 2 (placebo-treated), group 3 (MMSC injections), and group 4 (MF injections). Electrocardiography (ECG) and echo-cardiography were carried out in all animals before surgery, during 1 mo and 1 year after surgery. Myocardial perfusion rates were assessed with SPECT technique. IM dimensions (perfusion areas) were determined by means of staining heart sections with 2,3,5-TTC. Routine histology of myocardial sections was also performed. Vascularization rates of different myocardial areas were assessed in similar way.

Results

Intramyocardial transplantation of autologous bone marrow cells (MF and MMSC) into the area of rabbit experimental MI during the acute phase has changed natural process development and resulted in principally different morpho-functional outcomes. Ten days after coronary occlusion, all animals exhibited pronounced perfusion decrease in infarcted anterior wall of LV (p<0.05). However, the animals, treated with MMSC or MF showed less impaired perfusion rates than in control or placebo groups (p<0.05). 1.5 months after treatment, a gradual recovery of mean perfusion rates was observed in damaged myocardium in MMSC and MF-treated groups. These changes were confirmed by perfusion tomoscintigraphy.
Conclusions

MMSC injections were associated with pronounced therapeutic effect, by reducing myocardium damage area. By the contrary, MF transplantation showed a negative effect, expanding myocardium damage area and impairing systolic function indices. Both MMSC and MF intramyocardial transplantation display neoangiogenesis stimulation and perfusion improvement in rabbit experimental infarction area.

Keywords

myocardial infarction, experimental, bone marrow, mononuclear cells, mesenchymal stem cells, therapeutic injections, myocardial functions, histology.

Introduction

Myocardial infarction (MI) is a common reason of death and disability worldwide. Over the last decade, different cell therapy options have been tested to improve efficiency of MI treatment, including the s.c. cardiomyoplastics. Various cells are proposed to be used for this purpose. Autologous bone marrow is a widely used source of therapeutic cell preparations, thus providing freshly isolated mononuclear cell fraction (MF), or cultured multipotent mesenchimal stromal cells (MMSC). Meanwhile, existing works concerning clinical effects of MF and MMSC transplantation in acute MI yield controversial results [6, 11, 16, 24, 26, 29, 31], thus necessitating further studies and getting experience in this area.

The aim of the study

The aim of our experimental study was to compare the effects of intramyocardial MF and MMSC transplantation upon clinical course and outcome of MI, using a rabbit model during prolonged observation terms by using complex of modern functional and morphological study.

Materials and methods

The experiments were performed with male Chinchilla rabbits of 2.8± 0.2 kg weight, aged 3 - 4 months. All experimental procedures have been carried out in accordance with guidelines of local Ethical Committee at the First St.Petersburg I.Pavlov State Medical University.

Isolation, characterization, and morphological description of bone marrow cells

Bone marrow aspirate (10± 1mL) was obtained from a rabbit by iliac puncture, following premedication with Droperidol (0.5 mg/kg, Xylasine, 14 mg/kg body mass), and local anesthesia with 0.5 per cent Novocaine, and placed into a tube containing CPDS (citrate phosphate dextrose solution, Terumo, Japan). Mononuclear cell fraction was obtained by means of centrifugation (1600 g, 20 min) Percoll density gradient (63%). Interphase cell fraction containing nucleated cells was washed in Ca²⁺ and Mg²⁺-free Hanks’ solution (Gibco, USA), and span down by centrifuging. MMSC cell culture was obtained by MF cell passaging in α-MEM medium (ICN, USA) with 10 per cent fetal calf serum (HyClone, New Zealand) supplied with gentamycin sulfate (50 mcg/mL; Invitrogen, Great Britain) in a CO₂ incubator, at 5 per cent CO₂ and 100% humidity for three weeks. The medium was changed twice a week. Ascorbic acid (ICN, USA) was added at a final concentration of 50 mcg/mL after first medium change. After reaching a semi-confluent state, the cells were reinoculated by means of 0.25% trypsin solution (Gibco, USA), and EDTA (0.02%, Gibco, USA). Before being transplanted, the cultured cells did not display any signs of spontaneous osteogenic differentiation, as evidenced by negative staining for alkaline phosphatase with a standard BCIP-NBT reagent (5-bromine-4-chloride-3-indolyl phosphate/nitroblue tetrazolium, Sigma, USA), or any features of adipocytic transformation detectable with Sudan III/IV mixture (BDH Chemicals Ltd, Great Britain). Meanwhile, special studies with specific induction of differentiation to osteogenic and adipocytic lineage have shown their multipotency, i.e., the cultivated cells had typical MMSC characteristics.
In some animals, MNC or MMSC were stained prior to transplant with Hoechst nuclear fluorescent dye (Sigma, USA, final concentration of 1 mcg/mL), by shaking for 60 min in a \( \text{CO}_2 \) incubator.

Nucleated marrow cells were counted in Buerker chamber, and their viability was assessed with Trypan Blue solution (Labtech, Russia). The cells labelled with Hoechst dye were detected by their blue nuclear fluorescence upon microscopy, using an «Axioskope» (Zeiss, Germany).

**Experimental modelling and treatment of myocardial infarction**

The rabbits were subjected to a left-sided thoracotomy under mechanical ventilation of lungs, followed by ligation of anterior descendent left coronary artery at a distance of 1 cm from the heart apex (Fig. 1, A, B). Ten minutes after the coronary occlusion, intramyocardial injections of ME or MMSC suspensions were performed into 6 points of the presumed infarction area, using insulin syringes, at a mean cell number of \( 2\pm0.2 \times 10^6 \) in 0.4 mL of a-MEM growth medium (modified Eagle medium), or with equivalent volumes of culture medium (placebo treatment), as shown in Fig. 1 C, D. Control animals were not subjected to myocardial injections. Surgical wounds were closed in layers, pneumothorax being eliminated by means of active air aspiration from the pleural cavity. The surviving animals were classified in four groups, each consisting of 13 animals, i.e., Group 1, (controls), Group 2 (placebo-treated), Group 3 (MMSC injections), and Group 4 (MF injections). Subgroups of ten animals were observed for 1 year after the surgery.

**Detection of labelled cells in the myocardium**

Twenty days after the surgical intervention, the subgroups of three rabbits from Groups 3 and 4 (subjected to myocardial injections) were sacrificed (this term was limited by the life-span of detectable fluorochrome in the cells). The hearts were teased in two parts (basal and apical), in order to get sections at the level of arterial ligature perpendicular to long axis of the heart. Apical part of the heart was pulsed and treated with trypsin-collagenase enzyme mixture (Sigma, USA). The dissociated cells were collected by centrifugation, placed onto the microscopic slide, and the proportion of labelled cells have been counted.

**Electrocardiography (ECG)**

ECG was carried out in all animals before surgery, at the days +3, +7, +30, and one year after surgery. The results were evaluated by the 2nd standard lead on an ECG.

**Heart ultrasonography**

EchoCG was performed with a Sequoia 512 echocardiograph (Acuson, USA), using a linear 13 MGz transducer, according to a standard detection technique. EchoCG was performed in all cases before the surgery, at day +14, and 1 year after the surgery. The procedure was carried out in parallel with ECG recording. The end-diastolic size of left ventricle was registered in M-scanning regimen (EDS, mm). Left ventricular systolic function was registered by means of 2D scanning, from the apical access, in four-chamber and two-chamber positions. EF percentage calculations were made by means of a modified Simpson disc method, using the built-in software of the sonographic device. Aortal blood flow velocity was evaluated at the level of aortal valve (\( \text{V}_{\text{Ao}} \), m/sec), using impulse-wave Doppler sonography. Dynamic features of the left ventricular wall were also studied in intervention/infarction area, e.g., dyskinesia, hypokinesia, akinesia, calcification, or ossification in the areas of transplantation.

**Myocardial perfusion evaluation**

Perfusion rates were assessed with SPECT technique, using radiopharmaceutical tracer (RP) Myoview (Nycomed, Great Britain) labelled with Tc-99m. RP was injected, at a single dose of 30-50 MBq, into marginal ear vein via a peripheral catheter. 10 to 15 min after injection, a SPECT evaluation was performed by double-detector gamma chamber E.Cam. var (Siemens, Germany). To get quantitative values for myocardial perfusion rates, a ratio of mean RP accumulation in damaged versus reference areas was calculated in every case, as arbitrary units (Fig. 2). Uniformity of perfusion was determined as a minimum-to-maximum ratios (in pixels) for damaged and reference areas. The study was performed before surgical intervention, at day +10, and at 1.5, 6, and 12 months after surgery in all groups of animals.
Sizing of infarcted area, LV dilatation measurements, histological studies

Dimensions of an experimental IM were determined by means of 2,3,5-triphenyltetrazolium chloride (TTC) staining of the heart sections, thus allowing to discern irreversibly damaged myocardial tissue from viable myocardium. Moreover, a routine histological evaluation was performed. All the animals surviving for 12 months post-surgery were subjected to euthanasia, followed by immediate heart extraction. The organ was rinsed in physiological saline and sectioned transversally with a special device below the ligature level into three segments of equal thickness, i.e., apical (1), middle (2), and basal sections (3), as shown in Fig. 3A.

Quantitative evaluation of vascularization in damaged myocardial area

We have examined the borderline areas adjacent to the myocardial scars. Each tissue specimen was routinely stained by Mallory, and five consequent microscopic fields were evaluated at a 400x magnification. We performed separate counts of regulated-type vessels, i.e., arterioles, capillaries, venules, as well as non-regulated type-sinusoids, followed by calculating a mean blood vessel number per microscopic field.

Safety evaluation of approaches for angiogenesis stimulation in ischemic myocardium

To assess safety of cell transplantation in the studied groups of animals, we compared some general parameters, e.g., intra- and post-surgical mortality, incidence of arrhythmia and septic/inflammatory complications early after surgery. Condition of heart and internal organs was macroscopically evaluated by pathoanathomical obduction data, looking for occurrence of neoplastic processes, and microscopically evaluated for presence of local pathological changes in the areas of cell injections, in particular, atypically differentiated cells, focal sclerosis, or osteogenesis.

Statistical processing

We used SPSS software for statistical data processing. With small number of observations, the significance of differences was determined by a non-parametric Wilcoxon-Mann-Whitney criterion. All the data were presented as means ± SD. The differences by P values of <0.05 were regarded as significant.
Results
Detection of labeled cells in damaged myocardium

Twenty days after treatment, both MFs and MMSCs were detectable as fluorescent cells, at, resp., 13±2% and 11±3% of initial numbers (Fig. 4).

Figure 4. ECG dynamics in studied groups

ECG dynamics in different experimental groups

All the animals subjected to experimental coronary occlusion, developed typical ECG features of acute MI with distinct typical dynamics. However, all animals from control (placebo-treated), or MF-transplanted groups exhibited ECG signs of transmural MI, whereas MMSCs-transplant animals displayed only subendocardial myocardial damage (Fig. 5). Heart rate disturbances (atrial extrasystoles) in early postoperative period were developing with similar frequency (65%) in all the groups under study, except for MMSC-transplanted animals that showed them 3 times less often (20%). Twelve months after surgery, ECG patterns in MMSC-treated animals returned to the initial state, as compared with scar changes in ECG observed in MF-treated animals (Fig. 5).

Initial EchoCG parameters did not differ between the groups under study. After the surgical intervention (coronary occlusion), a significant increase of EDSLV was observed, along with diminished EF, Vao. However, these disruptions were dissimilar for different groups, i.e., minimal for MMSC-treated animals and maximal for MNC-treated animals (Table 1, Fig. 6, 7). One year after surgical intervention, a more expressed positive effect of MMSC transplantation was observed, i.e., nearly full recovery of all cardiological parameters (p<0.05), absence of akinetic areas, that sufficiently differed from appropriate parameters in other groups. In controls and placebo group, the disturbances were long-standing, and a small akinetic area was formed in the apical segment. The most negative changes were detected among animals transplanted with MNC, which revealed drastic decrease in LV systolic function, increased LV size, development of extended akinetic areas, as compared with group 3, controls, and placebo group (p<0.05) (Table 1, Fig. 6, 7).
Abbreviations: LVEF, LV ejection fraction; EDSLV, end LV diastolic size; V Ao, blood flow velocity in ascending aorta; *, differences from control group are significant by p<0.05.

Table 1. Dynamics of LV systolic parameters from EchoCG data (M±m)

<table>
<thead>
<tr>
<th>Groups</th>
<th>Parameters and terms of the study</th>
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<tr>
<td></td>
<td>EF, %</td>
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<tr>
<td></td>
<td>Initial values 14 days after surgery 12 months after surgery</td>
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<tr>
<td></td>
<td>EDSLV, mm 14 days after surgery 12 months after surgery</td>
</tr>
<tr>
<td></td>
<td>V Ao, m/sec Initial values 14 days after surgery 12 months after surgery</td>
</tr>
<tr>
<td>№1 (controls)</td>
<td>62.2±4.6 45.8±6.1 46.5±6.4 14.4±1.0 17.0±1.4 16.9±0.8 95±4 57±6 71±4</td>
</tr>
<tr>
<td>№2 (placebo)</td>
<td>64.9±4.3 46.7±3.5 48.7±3.9 14.1±0.4 16.8±1.2 16.6±0.8 94±3 58±6 73±6</td>
</tr>
<tr>
<td>№3 (MMSC)</td>
<td>61.7±4.0 53.1±4.8 58.2±3.1* 14.7±0.9 15.2±0.7 14.9±1.2 96±4 81±6 89±4*</td>
</tr>
<tr>
<td>№4 (MNC)</td>
<td>59.7±3.2 38.8±2.9 28.7±4.1 14.9±1.2 17.9±0.8 19.8±0.9 97±5 55±4 45±3*</td>
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Dynamics of myocardial perfusion in the groups under study.

Cumulative findings concerning dynamics of myocardial perfusion in damaged areas are presented in Table 2. Before surgery, the perfusion parameters in all groups were within reference ranges. Ten days after coronary occlusion, all animals exhibited pronounced perfusion decrease in infarcted anterior wall of LV (p<0.05). However, this decrease varied in different groups of animals, i.e., animals treated with MMSC or MNC showed less impaired perfusion rates than in control or placebo groups (p<0.05). At 1.5 months after treatment, a gradual recovery of mean perfusion rates was observed in damaged area of myocardium in groups 3 and 4, reaching the pre-surgical rates by 3 months, in group 3 to a greater extent, without subsequent dynamics. In controls and placebo-treated group, the altered perfusion rates remained at similar levels at all the terms of observation.

Figure 6. Transverse sections (1, apical; 2, median; 3, basal) from infarcted area of the heart wall in different experimental groups (Mallory staining) 12 months after MI modeling/treatment.

Figure 7. Numbers of regulated vs. sinusoid-type microvessels per a space unit of MI area 12 months after modelling and treatment in compared groups of experimental animals. Ordinate, number of microvessels per a IM space unit; abscissa, groups under study.
Morphometric heart studies

12 months after experimental MI modeling/treatment, general macroscopic appearance of rabbit hearts was similar for control and placebo-treated groups, i.e., enlarged heart size, as compared to healthy hearts (intact animals), along with apical aneurism of LV. Distinct scarring of LV anterior or wall was evident for the hearts from MMSC-treated animals, however, without evolving aneurysm, and without sufficient increase of the heart size. Heart dimensions among MF-treated animals did significantly exceed normal size, and appropriate heart parameters of healthy rabbits, and the animals from groups 1, 2, and 3. The animals transplanted with MF had extensive aneurysms at the apex and anterior wall of LV (Fig. 9, 10, 11). Scarring area and LV dilatation index for control group were, resp., 20.2±1.9%, and 0.19±0.02; 20.8±1.6% and 0.18±0.03 for placebo group; 35.8±1.6% and 0.18±0.03 for control group were, resp., 20.2±1.9%, and 0.19±0.02; 35.8±1.6% and 0.18±0.03 for placebo group; 12 months after experimental MI modeling/treatment, both groups of animals. After MMSC transplantation, a particularly different heart morphology as compared to control or placebo groups. After MMSC transplantation, a particularly different heart morphology as compared to control or placebo groups. The animals treated with MMSCs exhibited the best characteristics, i.e., 6.0±2.6% and 0.11±0.02 (Fig. 12).

Histological pattern of myocardium

It was shown that intramyocardial MMSC vs. MF transplantation resulted, after 12 months, in the development of principally different heart morphology as compared to control or placebo groups of animals. After MMSC transplantation, a pattern of microfocal cardiocclerosis was observed, followed by recovery of myocardial structure, whereas MF injections were followed by development of fibrotic aneurism affecting both left and right ventricles.

Quantitative myocardial vascularization

12 months after MI modeling/treatment, a significant difference was revealed in vascularization patterns of damaged myocardium in the studied groups, i.e., the total microvessel numbers per 1 microscopic field were, resp., 15±3, 13±3, 7±1 and 6±2 for groups treated with MMSCs, MFs, controls and placebo-treated animals. After intramyocardial autotransplantation of bone marrow, the microvessel density in the area was not only increased over controls or placebo groups (p<0.05), but it exhibited a distinct type of microvessels showing a regulated type-wall, whereas control group recovered with development of sinusoid-type microvessels.

Safety evaluation of neoangiogenesis stimulation in ischemic myocardium

Intra- and post-operative mortality in all the studied groups did not significantly differ (a mean of 10±3% и 10±4%). Incidence of septic and inflammatory complications in the intervention area was, in general, significantly lower (p<0.05) among intramyocardially cell-treated animals (1) than in control or placebo group (3). No differences were here noted between MF and MMSC-treated groups. At macroscopic examination one year after surgical intervention, both groups of cell-treated animals did not show any signs of malignant neoplasia. Local examination of myocardium at the locations of cell injections did not reveal any cells with atypical differentiation (osteogenic or chondrogenic tissues).

Discussion

Rabbit model of myocardial infarction induced by ligation of anterior descendent left coronary artery proved to be a convenient and demonstrative technique for evaluation of therapeutic angiogenesis, testing the effects of vascular growth factors and cellular therapy [14]. The results obtained in present study have shown that intramyocardial transplantation of autologous bone marrow cells (MF versus MMSC) into the MI area in rabbits during acute period may consid-

### Table 2. Dynamics of TTC accumulation in damaged myocardial area after treatment of acute MI in the groups under study by perfusion data (SPECT evaluation, M±m).

<table>
<thead>
<tr>
<th>Groups</th>
<th>TTC accumulation in damaged area (arb.un.), terms of study</th>
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<tr>
<td></td>
<td>Before surgery</td>
</tr>
<tr>
<td>Controls</td>
<td>1.01±0.01</td>
</tr>
<tr>
<td>Placebo</td>
<td>1.01±0.02</td>
</tr>
<tr>
<td>MMSC</td>
<td>0.98±0.03</td>
</tr>
<tr>
<td>MF</td>
<td>1.00±0.01</td>
</tr>
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</table>

Descriptions: arb.units = mean rates of TTC accumulation in damaged area/ mean values of TTC accumulation in reference area; * - significance of differences as compared to control group (p<0.05)
MMSC transplantation, as compared to controls and placebo, was associated with definite reduction of the myocardial damage, and later (12 months after MI) this group of animals exhibited normalization of heart systolic function, ECG, myocardial perfusion. Meanwhile, upon morphological examination, the infarcted area was represented by fine focal cardiac sclerosis, without aneurism formation. The detected changes might have been caused by paracrine effect of transplanted cells and, probably, by their differentiation into various myocardial structures. At the first phase of MI characterized by death of the cardiomyocytes, the transplanted MMSC are able to inhibit apoptosis and produce a cytoprotective effect, thus allowing cardiomyocytes and transplanted cells to survive the ischemic state [6,16,30]. Moreover, the detected changes might be associated with changing degree of inflammatory response, due to effect of the transplant upon the cytokine profile. In the course of inflammation caused by ischemic damage of myocardium, there are two cytokine groups: pro-inflammatory factors ([interleukin-1α, 1β, 6, 8, tumor necrosis factor-α, β and others]) mostly secreted by macrophages and neutrophils, and anti-inflammatory molecules ([interleukin-4, 10, transforming growth factor-β and others]), mainly released by lymphocytes, monocytes and macrophages. It was shown in vitro that bone marrow MMSC, along with anti-inflammatory cytokines ([interleukin-1α, 1β, 6, 7, 8, tumor necrosis factor-α, β]) produce a broad spectrum of anti-inflammatory cytokines ([interleukin-4, 10, transforming growth factor-β]), as well as specific receptors for interleukin-1α, 1β, 1, 3, 4, 6, 7, 8, tumor necrosis factor-α, β [22]. Moreover, it has been shown that the bone marrow MMSC secrete different chemokines: MCP-1, RANTES, MIP-1alpha [23]. Most probably, such cytokine profile may induce significant population changes, e.g., quantitative increase of macrophages, lymphocytes and monocytes that are able to secrete anti-inflammatory cytokines. This mechanism allows to reduce the degree of inflammatory response and the damaged area.

Progression of the small-focal myocardial infarction and normalization of systolic functions after BM MMSC transplantation, as shown in our study, can be also explained by potential ability of transplanted cells to differentiate into cardiomyocytes. However, an opportunity of BM MMSCs differentiation into cardiomyocytes remains problematic. Most researchers note the ability of bone marrow MMSC to acquire cardiomyocyte-like phenotypes after transplantation into the damaged myocardium, as shown by synthesis of α-actinin, troponin-T, tropomyosin. However, they have not revealed developing intercellular structures and/or contractile functions [20, 21, 27].

Thus, the changes in myocardial morphology and function after MMSC transplantation may be caused by their cytoprotective activity and ability to reduce the inflammatory response during acute infarction phase. By the contrary, intramyocardial MF transplantation, as compared to control and placebo groups, caused alterations of myocardial contractile function, expanding damage area, resulting into significant left ventricular fibroid aneurysms. Such effect is probably determined by MF ability to intensify inflammatory response, due to changing quantitative and qualitative cytokine profile in the damaged area, following direct massive injections of leukocytes. The bone marrow MFs constitute a heterogeneous population, containing mesenchymal stromal cells, hematopoietic stem cells, progenitor endothelial cells, as well as myeloid and lymphoid cells at different stages of maturation. There is an abundance of mature leukocytes among the bone marrow MF. Apparently, upon entering the ischemic myocardium, the MF start to release a variety of proteolytic enzymes and anti-inflammatory cytokines which attract blood cells to the altered areas, thus intensifying the inflammatory response [8]. The key point of myocardial repair is based on equilibrium between extracellular matrix degradation (needed for cell migration), and its resynthesis by the cells migrating to the damaged area. The proteolytic enzymes released by MF lyse intercellular collagen links and activate matrix metalloproteinases. Extracellular matrix degradation and expansion of infarcted area are caused by a cascade of proteolytic reactions. Moreover, the matrix metalloproteinases are also activated by some cytokines, e.g., as interleukin-1β in tumor necrosis factor-α, β [9, 19].

G. Ertl, S. Frantz have shown that dilatation of left ventricle and mortality were less expressed in the mice deficient for interleukin-1β than in animals with normal cytokine levels [7]. In addition, proteolytic enzymes released by transplanted MF of bone marrow induce massive death of cardiomyocytes. Cardiac myosin released from necrotized myocardium is a proven potent autoantigen. The autoimmune reactions occur via activation of T lymphocytes followed by the secondary T-cell induced damage of cardiomyocytes and altered remodeling of extracellular matrix [2, 28].

Thus, morphological and functional changes revealed in myocardium after bone marrow MF injections may be caused by excessive intensification of inflammatory response during the hyperacute phase of MI.

The results of our study indicate that intramyocardial transplantation of MMSC and MF of bone marrow leads to increased vascularization of damage area. We assume that the angiogenic effect of these cells is accomplished by secretion of vascular growth factors and, probably, by differentiation of transplanted cells into cellular components of the vascular wall.

MMSCs are able to secrete angiogenic factors, such as VEGF (vascular endothelial growth factor), bFGF (basic fibroblast growth factor), HGF (hepatocyte growth factor), TGF-β (transforming growth factor), angiopoietin-1 [13, 19]. Tang et al. (2005) demonstrated that the BM MMSC transplantation into the ischemic myocardium results into secretion of cytokine SDF-1 (stromal cell derived factor-1), which is a chemoattractant for endothelial cell progenitors and hematopoietic stem cells [22]. Progenitor endothelial cells migrating from peripheral blood into the damaged area are able to transform into mature endothelial cells and participate in angiogenesis [4, 12, 17]. Another mechanism of MMSCs involvement in post-transplant neoangiogenesis in myocardium is connected with their ability to differentiate into
endothelial cells, smooth muscle cells, pericytes, fibroblasts that subsequently become components of the vascular wall, as shown in several studies [4,22].

Involvement of bone marrow MF in angiogenesis, according to the literature data, is regulated by several mechanisms. First of all, the MF populations, e.g., mesenchymal stromal cells, hematopoietic stem cells, progenitor endothelial cells, as well as immature cells of myeloid and lymphoid series, are secreting various angiogenic factors (VEGF, bFGF, TGF-β, PDGF, angiopoietin-1), which stimulate proliferation of vascular endothelium and endothelial progenitors of damaged myocardium [13,25, 30, 31]. Secondly, the MF cells are able to secrete SDF-1 (stromal cell derived factor-1), which is a chemotactic agent for progenitor endothelial cells and hematopoietic stem cells [20]. Thirdly, such mesenchymal stromal cells within the MF fraction are able to differentiate into endothelial cells and other vascular structures [22].

Our data differ from other studies that did not show such significant differences after MF and MMSC transplantation to the experimental MI area [6,13,16,21,25,27,31]. A probable explanation is that transplantation of bone marrow cells in these studies was performed either in later terms following MI, or the cells were administered by a different route, e.g., via coronary arteries, or intravenously [1,3,5,10,24,29]. A prolonged observation in our study did not register such complications as ossification or formation of angiomatosis in the cell transplantation area, as previously described [15], neither any neoplastic processes have been registered.

**Conclusions**

1. Local intramyocardial transplantation of autologous bone marrow cells (MF and MMSC) into the area of rabbit, when performed in acute phase of experimental MI, was accompanied by changes in the MI development and resulted into different morphofunctional outcomes. I.e., transplantation of mesenchymal stem cells exerted a pronounced therapeutic effect, causing reduction of myocardial damage area and improvement of systolic functional indices. On the contrary, mononuclear cell transplantation produced negative effects, causing expansion of the damaged area and impairment of systolic indices.

2. Both MMSC and MF intramyocardial transplantation display stimulation of neoangiogenesis and improvement of perfusion in the infarcted area.

**Conflict of interest**

All the authors have no conflict of interest to declare.

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Сравнительные эффекты внутримиокардиальной аутотрансплантации различных видов клеток костного мозга на исходы экспериментального инфаркта миокарда у кроликов

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Резюме


24. van der Spoel TI, Jansen of Lorkeers SJ, Agostoni P et al. Human relevance of pre-clinical studies in stem cell therapy:


