THE CAUSE OF REPRODUCTIVE DISORDERS IN WOMEN FREQUENTLY IS THE INFLAMMATION OF PELVIC ORGANS. CHRONIC INFLAMMATION OF THE OVARIAS (OOPHORITIS) IS A COMMON DISEASE, ACCOMPANIED BY VARIOUS GYNECOLOGICAL DISORDERS, AND OCCURS INDEPENDENTLY AS WELL. ACCORDING TO THE MINISTRY OF HEALTHCARE OF UKRAINE, THE FREQUENCY OF INFLAMMATORY DISEASES OF PELvic organs IN THE STRUCTURE OF PATHOLOGY IN GYNECOLOGICAL PRACTICE REACHES 60–65% [1]. AS A CONSEQUENCE, PROLONGED AND RECURRENCE COURSE OF THE DISEASE IN 60% OF CASES CAN LEAD TO INFERTILITY. IT SHOULD BE NOTED THAT OOPHORITIS LARGELY OCCURS DUE TO THE PRESENCE OF PRIMARY IMMUNODEFICIENCY, WHICH FACILITATES THE TRANSITION OF ACUTE INFLAMMATORY PROCESS AT A CHRONIC STAGE. AT THE SAME TIME PROLONGED INFLAMMATION CAUSES A DECREASE IN IMMUNOLOGICAL REACTIVITY OF AN ORGANISM. IN CHRONIC INFLAMMATORY PROCESSES THE CHANGE OF CYTOKINE PROFILE IS NATURAL, WHICH LEADS TO STATISTICALLY SIGNIFICANT DECLINE IN THE EFFICIENCY OF INFERTILITY TREATMENT [2].

DURING THE TREATMENT OF UTERINE APPENDAGES’ INFLAMMATION THE PATIENTS RECEIVE MEDICINES OF ALMOST ALL PHARMACOLOGICAL GROUPS, INCLUDING THE RESERVE ANTIBiotics AND NON-Steroidal ANTI-INFLAMMATORY DRUGS THAT LEAD TO FORMATION OF ANTIBIOTIC-RESISTANT STRAINS, ACTIVATION OF OPPORTUNISTIC FLORA, INCREASED ALLERGIZATION, DYSFUNCTION OF THE IMMUNE SYSTEM [1, 2]. AS A RULE THE EFFECTIVENESS OF OOPHORITIS TREATMENT IS LOW, AND THEREFORE THE RESEARCHES OF ALTERNATIVE METHODS OF TREATMENT ARE INTENSIVELY CARRIED OUT, PARTICULARLY THE USE OF CELL THERAPY. TAKING INTO ACCOUNT THE ABOVEMENTIONED IT IS REASONABLE TO SEARCH FOR NEW MEDICINES HAVING THE ABILITY TO IMPACT ON THE CHRONIC PHASE...
of inflammation. The analysis of available reports indicates the relevance of research
purposely to explore opportunities to improve
reproductive function in animals and humans
using cell-tissue therapy. For example, the
cryopreserved placenta extract is used as a
stimulant of reproductive function [3]. The
question about the effectiveness of medicines,
which include the cryopreserved components,
such as stem cells or biologically active
components, is timely and appropriate one for
solving the task of correction of hormonal and
reproductive dysfunction.

There is an evidence of positive impact
of multipotent mesenchymal stromal cells
(MMSCs) in treatment of gynecological
diseases [4–7]. Carrying out the research, we
used bone marrow derived MMSCs. It is known
that MMSCs are identified by their ability to
adhesion in vitro, differentiation into various
types of tissues and the expression of several
cell surface markers, among which the most
typical are CD 105, CD 73, CD 90, at the
absence of CD 34, CD 45 markers [8]. Modern
technologies of culture and cryopreservation
make possible receiving the stock of autologous
cell with a subsequent long-term storage
at low temperatures without significant
changes in functional status.

Existing data about the introduction of
MMSCs into intact animals indicate immune
modulating effect and diffuse distribution of
donor’s material in the tissues of mesoderm
origin [6, 9]. For assessment the impact of
transplanted cellular material to restore
the damaged tissue or organ histological,
biochemical, biophysical, immunological
methods are used, however, the location of
the introduced cells is not always determined.
According to this a simultaneous assessment of
the ability of cryopreserved MMSCs to homing
and stimulation of regenerative processes is
important. This allows determining the term of
a cell keeping in a tissue being restored, and the
regenerative potential of the cells themselves.

The aim of this work was to investigate the
effect of cryopreserved bone marrow derived
multipotent mesenchymal stromal cells on
reparative regeneration and to determine
their localization under intravenously
administration in the animals with chronic
inflammation of the ovaries.

**Materials and Methods**

MMSCs of the femur bone marrow of mice
\((n = 5)\) was used in the study. The cells were
isolated by washing out with Hanks solution
(PAA, Austria) followed by flushing through
a needle with gradually decreased diameter.
The next step was centrifugation at 1500 rpm
\((834 \text{ g})\) for 5 min. The cell suspension was
resuspended in culture medium and plated on
a culture flask (PAA) with \(10^5\) cells per \(\text{cm}^2\)
density. Cultural medium contained: Iskove’s
Modified Dulbecco’s Mediume (PAA), 10% fctal
bovine serum (FBS) (HyClone, USA),
gentamicin (150 mg / ml) (Farmak, Ukraine)
and amphotericin B (10 mg / ml) (PAA).
Cultural medium was changed every 3 days.
In the study we used standard conditions of
culture at \(37^\circ\text{C}\) in the atmosphere of 5% \(\text{CO}_2\).
Cell culture was passaged after reaching a
confluent monolayer.

Cell cryopreservation was carried out in
in culture medium supplemented with
10% DMSO (PAA) and 20% FBS. 1 ml of
 suspension was placed in cryovials (Nunc,
USA). Cryopreservation was performed with
a programmable freezer ZPM-1 (SDTB with
PP IPC&C NAS of Ukraine). Cooling rate was
1 deg/min down to \(-80^\circ\text{C}\), with following
transfer into liquid nitrogen [8]. Samples
were stored under the low temperature bank
conditions for 4 months, thawed on water bath
at \(40^\circ\text{C}\) up to the liquid phase. Removing of
cryoprotectant was performed by slow addition
of 1:9 Hanks solution (PAA), followed by
centrifugation at 1500 rpm(834 g) for 5 min.
Cell viability was assessed by exclusion of supravital
trypan blue dye (Sigma-Aldrich, USA).

As experimental objects we used 50 outbred
mature female white mice weighing 18–20 g.
In order to cause chronic inflammation of
the ovaries an inactivated vaccine of
*Staphylococcus aureus* strain 209 was
once injected intraperitoneal with insulin syringe
and then mice were kept in the animal house
conditions for 21 days without treatment.
For vaccine we prepared a daily culture of
*Staphylococcus aureus* by the standard method
[10]. The resulted culture was washed out
from agar with 5 ml of saline solution with
titer determination by the method of standard
dilutions and inactivated by incubation at \(75^\circ\text{C}\)
during 1h. After incubation the suspension
of inactivated bacterial cells was diluted to
\(50\times10^6\) microbes’ bodies in 0.3 ml of saline
solution per animal.

To the 22\text{nd} day after the administration
of inactivated vaccine the control and
experimental groups were formed when in
the tail vein the following were injected:
control group \((n = 15)\) — saline solution
\((0.2 \text{ ml})\); experimental group \((n = 15)\) —
cryopreserved bone marrow derived MMSCs
(CrMMSCs); experimental group 2 \((n = 10)\) — bone marrow derived CrMMSCs labeled PKH-26. The volume of administrated liquid in all the experimental groups was 0.2 ml per animal \((0.5 \times 10^5\) viable CrMMSCs); control group 2 — intact animals \((n = 10)\) of similar weight and age without manipulation. Insertion of PKH-26 (Sigma-Aldrich) in CrMMSCs was carried out according to the instructions of the manufacturer.

The animals were removed from the experiment to the 10th and 21st day after cell therapy by dislocation of the cervical vertebrae. For histological study and detection of entered cell location, ovaries were cut off with the surrounding tissue of omentum. Material for histological study was fixed in the 10\% aqueous neutral formalin and then the serial paraffin sections with 4–5 µm thickness were done and stained with hematoxylin-eosin. The number of follicles and their development stages were classified as reported [11], modified by Gougeon [12] with the microscope.

To assess the state of the oocytes they were mechanically obtained from the ovaries of animals of the control groups and experimental group 1 to the 21st day of therapy. Presence/absence of apoptosis in the oocytes was assessed with Annexin-V (Becton-Dickinson, USA). Staining was performed according to the standard procedure of the manufacturer. Oocyte microimages were performed with the fluorescent microscope (MIKMED-2, Russia).

To investigate the labeled cell localization in the animal organism, the ovaries were isolated and the cryostat sections (7 µm) were prepared. The cryostat samples were investigated with fluorescent microscope. Autofluorescence was blocked with 0.3M glycine solution (PAA) by 20-min incubation and followed by microscopy [13]. For detection of labeled PKH-26 MMSCs frozen slices were additionally stained with 1 µg/ml DAPI (Sigma-Aldrich) for nuclei visualization. Results were fixed by photographing.

All the manipulations with the animals were carried out in accordance to the requirements of the “European Convention for the Protection of Vertebrate Animals used for Experimental and other Scientific Purposes” (Strasbourg, 1986), “General Principles of Animal Experiments”, approved by II National Congress of Bioethics (Kyiv, 2004), and the rules of the Commission of Bioethics of the IPC&C NAS of Ukraine.

In statistical analysis we used a single-factor analysis of variance and Student’s t-test using Statistica 8 software.

Results and Discussion

At the first stage the viability of MMSCs were assessed by trypan blue staining, the results were \(95.2 \pm 4.6\%\) and \(78.5 \pm 6.2\%\) for native and cryopreserved MMSCs, respectively.

The results of macroscopic research of an abdominal cavity showed that animals of the control and experimental groups had a normal blood supply in the vessels, a moderate amount of fat around the ovaries without visual signs of inflammatory reaction and exudation at all stages of observation.

Histological structure of intact animals’ ovaries corresponded to the standards of mature mice with no signs of inflammatory changes (Fig. 1). Ovarian surface was covered with a single layer of cubical epithelium, under which tunica albuginea was located. Follicular structure of the samples was presented by primordial, primary, preantral and antral follicles that were separated by a stroma. The environment of antral follicles was homogeneous with weak eosin staining; oocytes were without signs of degeneration with evenly surrounded granulosa cells. Single atretic follicles with typical histological structure were defined on the periphery of sections. Corpora lutea were filled with radial strands of luteum cells.

In the central part of histological sections of the ovaries of animals from group 1 with chronic inflammation after injection of saline solution the stromal fibroblast-like cells which were surrounded by single developing follicles were observed to the 10th day (Fig. 2, A). The leukocyte infiltration was observed at the edges and in the center of all the samples. It should be noted a significant amount of atretic follicles, which were characterized by the contour deformation. Atresia occurred according to the

Fig. 1. Histology section of intact animal ovary. Here and in Fig. 2, 3: hematoxylin and eosin staining. ×250
type of productive process with a multilayer membrana granulosa forming. Single primary follicles with the signs of degeneration were detected along the section periphery. In such follicles the layers of cuboidal granulosa cells were not only compressed but also separated and were less closely adjacent to an oocyte. It is known that disorder in a contact between oocyte and granulosa cells is an irreversible process, adversely affecting an oocyte metabolism and development [14].

Further degradation of the structural components of ovarian tissue was occurred with the increase in observation period to the 21\textsuperscript{th} day in the animals with oophoritis after injection of saline solution (Fig. 2, B). Thus, the area of ovarian section was filled with stromal interstitial cells. Primordial and primary follicles in most cases were not available, only a few preantral follicles with signs of degeneration were noted on the periphery of histological section.

Compared with the control group 1 the analysis of histological research of the ovaries of the animals with CrMMSC therapy showed that the morphological structure of the ovaries had a positive recovery dynamics to the 10\textsuperscript{th} day after the cell injection (Fig. 3, A). Along the periphery of histological sections of ovaries, amount increasing of follicles of both early and late stages of follicle genesis was observed. Follicular profile was presented by the primordial, primary and preantral stages of oocytes. The intensity of leukocyte infiltration in ovarian tissue was significantly lower compared with the control group 1.

The results of histological examination of the ovaries in animals with the oophoritis and CrMMSC therapy to the 21\textsuperscript{st} day (Fig. 3, B) showed a reparative regeneration with a tendency to normalization of morphological parameters of ovarian tissue. Slight leukocyte infiltration was observed on the surface of ovarian cortical layer that pointed to the reduction of inflammation intensity in experimental preparations relative to the control group 1. The follicular profile was characterized by the presence of primordial, primary, preantral and single antral follicles, which were located at the edge of sections. In the ovaries of the animals with bone marrow CrMMSC therapy as well as extinction inflammatory signs the activation of regenerative processes took place, while in the control animals with administered saline solution they were not observed.

The results of total number of follicles per ovary counting in histological sections (Fig. 4) to the 10\textsuperscript{th} day showed statistically significant reduction ($P < 0.05$) of this parameter both in the control and experimental groups up to the values of $10.5 \pm 2.48$ and $9.4 \pm 2.63$ respectively compared with intact animals ($18.3 \pm 4.52$). With the increasing of observation period up to the 21\textsuperscript{th} day, the number of follicles in the control group 1 continued decreasing ($7.4 \pm 2.18$), while in the experimental group this parameter significantly increased ($P < 0.05$) up to $15.3 \pm 1.8$, although it did not reach the values of the control group 2.

It is known that the oocyte quality affects significantly on the viability of embryos at pre-implantation stages of development, the probability of pregnancy and fetal development. Therefore, it is important to determine the presence/absence of apoptotic features in oocytes. These results are shown in Fig. 5. In the oocytes of the control group 2 the signs of apoptosis were not fixed. In the oocytes of the control group 1 together with the decrease of total number of oocytes $85.3 \pm 5.2\%$ ones had the apoptosis signs (Fig. 5, A, B). Number of oocytes with apoptotic signs was $5.7 \pm 0.8\%$ in the animals with CrMMSC therapy (Fig. 5, C, D). Thus, there was detected a tendency to restore the total number of oocytes in the ovaries of experimental animals after 3 weeks of intravenous injection of

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig2.jpg}
\caption{Histology sections of ovaries of animals with chronic inflammation and administration of saline: The 10\textsuperscript{th} (A) and 21\textsuperscript{st} (B) days}
\end{figure}
CrMMSCs, that can indicate the activation of folliculogenesis after cell therapy.

At the final stage we determined the localization of CrMMSCs after their intravenous injection into mice with oophoritis. It is known that the detection of transplanted cells in the recipient’s organism is performed with the help of histochemical, radioimmunological methods and fluorescent microscopy. For this purpose the nanoparticles, fluorescent dyes, embedded genetic fluorescent probes and other agents that promote detection of the introduced cell location are injected into the cells prior to their application. We have chosen the probe PKH-26, which according to the information of developer is a lipophilic dye, non-radioactive substance that binds to cell membranes and shows no toxic effects [15]. Fig. 6 shows micrographs of bone marrow MMSC culture labeled with PKH-26 dye. There was observed the presence of luminescence in a red region of the spectrum in the cryostat sections of ovaries to the 10th and 21st day after CrMMSC therapy. To the 10th day the emission intensity was weak and had the shape of small-sized conglomerates, diffusely located in the center of the samples (Fig. 7, A). To the 21st day the luminescent objects appeared and looked like a small sized conglomerates in the center and edge part of the sections (Fig. 7, B).

It should be noted that the number of luminescent objects in cryostat sections of ovaries was lower to the 21st day compared to the previous period. It is known that PKH-26 dye does not pass from a cell to cell, but has the ability to transition into daughter cells by mitosis. Thus, the luminescing cells in the examined tissues can be directly introduced cells as well as daughter ones.

The phenomenon of directed migration of stem cells to the area of injury, ischemic or neoplastic lesions was an important step in understanding the mechanisms of regeneration tissue processes. Due to homing, which existence was proven in many studies, there is a migration of cells to the damaged and ischemic areas [16–18] to implement the cell therapy effects at an organism level. To date, 79 cytokines, growth factors and hemoatractants and over 20 types of receptors have been identified to involve in the directed migration processes of various types of stem cells in a norm and pathology [19]. The central role in this process belongs to the interaction between SDF-1α factor (its content increases in the areas of injury) and receptor CXCR4 on the MMSCs [20, 21]; moreover there is discussed the importance of ligand-receptor interactions of SCF–Kit [22], HGF/c-met [23] and others.

The studies indicate that injected in the tail vein MMSCs have been determined in ovarian tissue to the 10th and 21st day after treatment. This is one more proof of the hypothesis.
that MMSCs are not only passively carried throughout the body by a blood flow, but also perform a directed migration to the area of inflammation of damaged tissues.

According to the date [24, 25], the bone marrow MMSCs have immune regulatory effect except their regenerative properties. Due to the last administrated cells indirectly influence the course of inflammation in the ovaries. The results of cryopreserved MMSC influence on the process of ovarian recovery allow concluding about their possible usage in therapy of oophoritis.

It should be noted that intravenous cell injection, including MMSCs, is considered to be safe and non-invasive method of therapy. For this reason the systemic MMSCs injection may be the most appropriate in the treatment of considered pathology. However, there remains open the question about more remote effects of cell therapy, including effects on fertility, reduced under chronic inflammation.

The results in animals suggest that intravenous MMSC injection can be effective for treatment of inflammation in human ovaries. Perhaps the mechanisms of directed migration and stem cells homing to the source of injury belong to the complex of tissue homeostasis system, which regulates the effector functions of damaged cells modulating the survival processes and apoptosis, proliferation and differentiation.

The development of regenerative cell therapy is the direction of medical biotechnology, which is rapidly growing and getting usage in the treatment of gynecological pathology [4–6]. Based on the results of this study, we pay attention of researchers to the fact that bone marrow MMSCs can be considered as a potential object of cell therapy of gynecological diseases. However, this issue requires further study during longer period of time. In addition, it is important to identify the fertility characteristics of control animals with oophoritis and animals with CrMMSC therapy alone and in combination with standard anti-inflammatory drugs.
Thus, it can be argued that intravenous injection of cryopreserved bone marrow multipotent mesenchymal stromal cells into mice with chronic inflammation of the ovaries produces a modulatory effect on the course of inflammation and restoration of folliculogenesis without generating apoptosis in oocytes, as well as that cryopreserved bone marrow multipotent mesenchymal stromal cells at intravenous injection are detected in the animal ovaries to the 10th and 21st day.

REFERENCES

1. Dubossarskaja Z. M., Miljanovskij A. I., Koljadenko V. G. Chronic inflammation of internal female genitals. Kyiv: Zdorov'ja. 2003, P. 115–118. (In Russian).
2. Drannik G. N. Clinical Immunology and Allergology. Moscow: OOO Med. inform. agentstvo, 2003. 604 p. (In Russian).
3. Grishchenko N. G., Klimenko N. A., Gorgol' N. I. Tatarko S. V Effect of placental cryoextract on chronic inflammation of the ovaries in mice. Medycyna s'ohodni i zavtra. 2010, N 2–3, P. 7–17. (In Russian).
4. Takehara Y., Yabuchi A., Ezeo K., Kuroda T. The restorative effects of adipose-derived mesenchymal stem cells on damaged ovarian function. Labor. Invest. 2013, 93(2), 181–193.
5. Fu X., He Y., Xie C., Liu W. Bone marrow mesenchymal stem cell transplantation improves ovarian function and structure in rats with chemotherapy-induced ovarian damage. Cytotherapy, 2008, 10 (4), 353–363.
6. Weina L., Qixuan X., Junwen Q. Effect of mesenchymal stem cell transplantation on immunological injury of the ovary in mice. J. South Med. Univ. 2011, 31 (5), 825–829.
7. Komarova S., Roth J., Alvarez R., Pereboeva L. Targeting of mesenchymal stem cells to ovarian tumors via an artificial receptor. J. Ovar. Res. 2010, 3 (12), 12–18.
8. Haack-Sorensen M., Bindslev L., Mortensen S., Friis T., Kastrup J. The influence of freezing and storage on the characteristics and functions of human mesenchymal stromal cells isolated for storage on the characteristics and functions of human mesenchymal stromal cells isolated for clinical use. Cytotherapy, 2007, 9 (4), 328–337.
9. Dittmar T., Entschladen F. Migratory properties of mesenchymal stem cells. Adv. Biochem. Eng. Biotechnol. 2013, V. 129, P. 117–136.
10. Egorov N. S. Practical work on Microbiology. Moscow: MGU, 1976, 307 p. (In Russian).
11. Oktay K., Newton H., Mullan J. Development of human primordial follicles to antral stages in SCID/hpg mice stimulated with follicle stimulating hormone. Hum. Reprod. 1998, 13 (5), 1133–1138.
12. Gougeon A. Dynamics of follicular growth in the human: a model from preliminary results. Hum. Reprod. 1986, 1 (2), 81–87.
13. Baschong W., Swetterlin R., Laeng R. H. Control of autofluorescence of archival formaldehyde-fixed, paraffin-embedded tissue in confocal laser scanning microscopy (CLSM). J. Histochem. Cytochem. 2001, 49 (12), 1565–1572.
14. Kiroshkha V.V., Medinec E.A., Tishhenko Ju.O., Bondarenko T. P. Dynamics of changes in morphology neonatal ovarian tissue during cold storage, depending on the composition of the incubation medium. Problemy kriobiologii. 2012, 22 (1), 61–70. (In Russian).
15. Johnsson C., Festin R., Tufveson G., Totterman T. H. Ex vivo PKH26-labelling of lymphocytes for studies of cell migration in vivo. Scand. J. Immunol. 1997, 45 (5), 511–514.
16. Le Blanc K., Mougiakakos D. Multipotent mesenchymal stromal cells and the innate immune system. Nat. Rev. Immunol. 2012, 12 (5), 383–396.
17. Karp J. M., Leng G. S. Mesenchymal stem cell homing: the devil is in the details. Stem Cells. 2009, 4 (3), 206–216.
18. Lee J. S., Hong J. M., Moon G. J., Lee P. H., Ahn Y. H., Bang O. Y. A long-term follow-up study of intravenous autologous mesenchymal stem cell transplantation in patients with ischemic stroke. Stem Cells. 2010, 28 (6), 1099–1106.
19. Horuk R. Chemokine receptors. Cytokine and growth factor review. 2001, 12 (4), 313–335.
20. Kucia M., Reca R., Miecz P., Wanzek J., Wojakowski W, Janowska-Wieczorek A., Ratajczak J., Ratajczak M. Z. Trafficking of normal stem cell and metastasis of cancer stem cells involve similar mechanisms: Pivotal role of the SDF-1-CXCR4 axis. Stem Cells. 2009, 23 (7), 879–894.
21. Bhakta S., Hong P., Koc O. The surface adhesion molecule CXCR4 stimulates mesenchymal stem cell migration to stromal cell-derived factor-1 in vitro but does not decrease apoptosis under serum deprivation. Cardiovasc. Revasc. Med. 2006, 7 (1), 19–24.
22. Erlandson A., Larsson J., Forsberg-Nilsson K. Stem cell factor is a chemoattractant and a survival factor for the CNC stem cell. Exp. Cell Res. 2004, 301 (2), 201–210.
23. Wondergem R., Ecay T. W., Mahieu F. HGF/ SF and menthol increase human glioblastoma cell calcium and migration. Biochem. Biophys. Res. Commun. 2008, 372 (1), 210–215.
24. Krampera M., Cosmi L., Angeli R., Tabera S., Perez-Simon J. A., Diez-Campelo M. The effect of mesenchymal stem cells on the viability, proliferation and differentiation of B-lymphocytes. Haematologica. 2008, 93 (9), 1301–1309.
МУЛЬТИПОТЕНТНИ МЕЗЕНХІМНІ СТРОМАЛЬНІ КЛІТИНИ КІСТОВОГО МОЗКУ В ТЕРАПІЇ ХРОНІЧНОГО ЗАПАЛЕННЯ ЯЄЧНИКІВ У МИШЕЙ

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Метою роботи було дослідити вплив на хронічне запалення яєчників мишей та визначити локалізацію мультипотентних мезенхімних стромальних клітин кісткового мозку за внутрішньовенного введення. Результати гістологічного дослідження засвідчили, що в експериментальних тварин в умовах клітинної терапії відбувалась активізація репаративного процесу з тенденцією до нормалізації морфологічних параметрів оваріальної тканини на фоні згасання запальньої проявлень. На 21-шу добу в контрольній групі з уведенням фізіологічного розчину загальна кількість фолікулів була знижена (7,4 ± 2,18%) порівняно з інтактними тваринами (18,3 ± 4,52%), а 85,3 ± 5,2% ооцитів мали ознаки апоптозу (Annexin V+). В експериментальній групі кількість фолікулів достовірно збільшувалась до величини 15,3 ± 1,8%, а кількість апоптозних ооцитів зменшувалась (5,7 ± 0,8%) порівняно з контролем. Люмінесцентна мікроскопія кріостатних зрізів яєчників тварин після терапії міченими РКН-26 клітинами дала змогу виявити дифузне розподілення люмінесцентних об’єктів, які мали вигляд невеликих за розміром конгломератів клітин. Установлено, що криоконсервовані мультипотентні мезенхімні стромальні клітини кісткового мозку за умов внутрішньовенного введення активно впливають на репаративні процеси, сприяючи нормалізації морфологічних параметрів оваріальної тканини.

Ключові слова: хронічне запалення яєчників, мультипотентні мезенхімні стромальні клітини кісткового мозку, клітинна терапія.

МУЛЬТИПОТЕНТНЫЕ МЕЗЕНХИМНЫЕ СТРОМАЛЬНЫЕ КЛЕТКИ КОСТНОГО МОЗГА В ТЕРАПИИ ХРОНИЧЕСКОГО ВОСПАЛЕННИЯ ЯЧНИКОВ У МЫШЕЙ

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Целью работы было исследование влияния на хроническое воспаление яичников мышей и определение локализации криоконсервированных мультипотентных мезенхимных стромальных клеток костного мозга при внутривенном введении. Результаты гистологического исследования свидетельствовали, что у экспериментальных животных в условиях клеточной терапии происходила активизация репаративных процессов с тенденцией к нормализации морфологических параметров овариальной ткани на фоне угасания воспалительных проявленияй. На 21-е сутки в контрольной группе с введением физиологического раствора общее количество фолликулов было снижено (7,4 ± 2,18%) относительно интактных животных (18,3 ± 4,52%), а 85,3 ± 5,2% ооцитов имели признаки апоптоза (Annexin V+). В экспериментальной группе количество фолликулов достоверно увеличивалось до величины 15,3 ± 1,8% а количество апоптозных ооцитов снизилось (5,7 ± 0,8%) по сравнению с контролем. Люминесцентная микроскопия криостатных срезов яичников животных после терапии меченными РКН-26 клетками показала наличие диффузного распределения люминесцирующих объектов, которые имели вид небольших по размеру конгломератов клеток. Установлено, что криоконсервированные мультипотентные мезенхимные стромальные клетки костного мозга при внутривенном введении животным с хроническим воспалением яичников оказывают модулирующее влияние на течение воспаления, способствуют восстановлению фолликулогенеза и определяются в яичниках экспериментальных животных на 10- и 21-е сутки после введения.

Ключевые слова: хроническое воспаление яичников, мультипотентные мезенхимные стромальные клетки костного мозга, клеточная терапия.