EXPERIMENTAL STUDY

Evaluation of the repair of diaphyseal fracture of femoral bone using bone marrow mesenchymal stem cells in nicotine-bearing rat

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Abstract

OBJECTIVE: Following bone fractures during accidents, some patients suffer from poor repair of bone fractures and subsequent aesthetic and psychological problems. One of the treatments is based on transplantation of stem cells (seeded on scaffolds) to the lesion site. Bone marrow stromal cells (BMSCs) are multivalent stem cells which are able to reproduce and differentiate into osteogenic cells. The objective of this study was to evaluate the treatment of bone fractures by means of transplantation of the latter cells in rats.

METHODS: In this study, the therapeutic effect of mesenchymal stem cells from bone marrow adipocytes was evaluated in bone fractures. BMSCs were isolated from rat femur. Two sources of differentiated and non-differentiated osteocyte cells were provided and mixed with collagen in order to be transferred to animals divided in three main groups of model: nicotine, non-nicotine and control groups. After four weeks, the repair of the fracture that had been inflicted by a 2-mm drill into the diaphyseal region of the femoral bone was investigated by radiographic tests and histopathologic staining.

RESULTS: The radiographic results as well as those of histopathologic staining showed that osteogenesis was more intensive in the non-nicotine group than in the nicotine group with differentiated and non-differentiated osteocyte cells.

CONCLUSION: The transplantation of differentiated BMSCs to a bone lesion affected the repair of bone fractures while the nicotine agent played an important role in delaying the bone regeneration (Tab. 1, Fig. 8, Ref. 31).

KEY WORDS: bone marrow mesenchymal stem cell, nicotine, rat, bone fracture regeneration.

Introduction

One of the most common abnormalities in bone fractures is the slow and imperfect repair. Bone density reduces with age and makes bones prone to fractures. Imperfect repair brings about disabilities, which incur high cost for both the individual and society. Due to the relative limitation of repair in the above conditions, researchers try to find a suitable treatment for these patients. There are several modes of therapy for quick repair of bone fractures. One of these modes is cell therapy. In order to gain the best outcome from cell transplantation, the cells required for this propose should have features such as easy access, rapid division in the culture medium, immunology that is neutral or synchronized with recipient tissue, capability of long-term survival and compatibility with host tissue. According to these characteristics, bone marrow stromal cells (BMSCs) seem to be suitable alternatives for this purpose (1, 2). Mesenchymal stem cells derived from bone marrow (BMSCs), which have the potential to form different types of mesenchymal tissues, are also the cellular source for bone tissue engineering applications. From the scientific point of view, the stem cell is considered to be an undifferentiated cell with the abilities of multiplication, self-renewal, and production of different cell lines. The bone marrow is a soft tissue full of blood vessels with scaffolds of reticular strata that blood cells, adipocytes and stem cells are placed on (3–5). In this tissue, the mesenchymal cells that are located in the bone marrow cavities are able to transform into a variety of other cells, including chondrocytes, osteoblasts, fibroblasts, adipocytes, endothelial cells and myocytes (6–8). The mesenchymal stem cells in the bone marrow have an extensive association with various cells such as blood vessel endothelial cells, osteoblasts, reticular cells of the red bone marrow and adipocytes (9, 10). This association is essential for processes such as hematopoiesis, production and maturation of defense cells, and regulation of immune response (11). It has also
been shown that the connection between mesenchymal stem cells and these cells is through both physical and chemical mechanisms such as production and secretion of molecules and growth factors (12). Under normal condition, the mesenchymal stem cells in the bone marrow are in a non-differentiated state (13). The repair mechanisms activated by stem cells include the proliferation and differentiation of damaged cells, control of the programmed cell death, mobilisation of other cells from other areas affecting the repair, and control and modification of immune response in the injured area (14–16). Although the mentioned mechanisms do not fully cover the repair in the tissue, these mechanisms can be considered to be the basis for damaged tissue repair. In general, there are two main reasons for the use of mesenchymal stem cells for the therapeutic purpose, including the creation and replacement of damaged cells and secretion of factors effective in the repair process performed by mesenchymal stem cells. The creation and replacement of damaged cells form the basis of cell therapy in restorative medicine. The secretion of factors effective in the repair process performed by mesenchymal stem cells is considered to be the pivotal element also in many cellular processes (17). The purpose of this study was to evaluate the repair effect of mesenchymal stem cells from rat bone marrow in rat diaphyseal bone fractures.

Materials and methods

Isolation and culture of mesenchymal stem cells of rat bone marrow

In order to isolate and culture the mesenchymal stem cells of bone marrow, Wistar male rats (Rattus norvegicus; 200 to 250 g) were purchased from a laboratory of experimental animals at Baqiyatallah University of Medical Sciences (Tehran, Iran). The animals were anesthetized by inhalation of carbon dioxide. The rats were immersed in 70 % alcohol, then in iodine and again in 70 % alcohol to eliminate the contaminating agents on the body surface. The femur and tibia were removed from both sides and the tissues adhering to the bones were cleaned off. Next, the cleaned bones were soaked in PBS containing antibiotics, namely penicillin (100 unit/ml) and streptomycin (100 mg/ml) and placed under laminar flow. Under sterile conditions, the DMEM-Low glucose culture medium was added to 10 ml syringes. The two ends of the bones were cut and the contents was drained by flushing method with syringes into 25-ml cell culture flasks. Subsequently, 10 % FBS, 1 % Pen/Strep and finally, low-glucose DMEM were added to reach volume of 5 ml. The flasks were incubated in 5 % CO₂ at 37 °C. The next day, the floating cells were washed with PBS and adherent cells were cultured for 14 days while changing the media every three days. Following 14 days, when the confluency of cells reached 80 %, the first sub-culture was performed using Trypsin/EDTA 2.5 % solution to allow cells to grow and proliferate. Briefly, the cell culture media was removed and the cells were washed with PBS. Then the cells were incubated for 3–5 min with 0.5 ml of 1X trypsin. After detaching the cells, the media containing FBS was added to the flask to neutralize the effect of trypsin. The cells were pelleted at 600 g for 8 min (1200 rpm for 7 min). The pellet was resuspended in 1 ml of new culture media. Ten microliters of the cell suspension were stained with 0.4 % trypan blue and counted using Neubar glass slide under the inverted microscope. The percentage of live cells and total number of cells were obtained. Considering the ability of population growth (doubling time), the cells were transferred to other flasks with the ratio of 1 to 2 or 1 to 3. The second passage of cells was performed in the same way.

Evaluation of cell surface specific markers

To evaluate the surface markers, the cells from the third sub-culture were used. The cells were detached using trypsin and resuspended in 1 ml PBS. Then, 50 μl of cell suspension containing 10⁶ cells was transferred to each tube of test and isotype control. The amount of 5 μl of the CD90, CD106, CD34 and CD45 monoclonal antibodies were added to the test tubes. The amount of 5 μl of IgG-FITC antibody against rat cells was added to the isotype control tube to identify and remove non-specific bonds. The tubes were placed in darkness for 30 min at 4 °C. Then, a volume of 500 μl PBS was added to the sample and washed. Cells were then fixed by 50 μl of paraformaldehyde solution (the paraformaldehyde solution can be added simultaneously with the antibody) and then the fluorescence of the samples was examined by flow cytometry (18).

Differentiation into osteocyte and adipocyte

In this study, the osteocyte differentiation test was used to verify the nature mesenchymal cells and their differentiation potency. The media of the third subculture of confluent cells was changed with bone differentiation media and the cell differentiation culture was maintained for 21 days. The osteocyte differentiation medium contained DMEM containing 50 μg/ml ascorbic acid 3-phosphate, 10 nm dexamethasone and 10 mM beta-glycerol phosphate. For differentiation into cartilage, 0.5 ml of chondrogenic media including DMEM containing 10 ng TGF-B3 growth factor, 50 mg ITS+premix, 10 ng BMP-6, 3.55 mg linoleic acid, 1.25 mg bovine serum albumin and 2 % bovine serum was added to the cell pellet at the bottom of the tube. Adipocyte differentiation was performed as follows-. The third sub-cultures of confluent cells were cultured in a 6-well plate. After covering the cells with adipogenic medium (DMEM), the media containing 10 % FBS, 50 μg/ml ascorbic acid 2-phosphate, 100 μm dexamethasone and 50 μg/ml indomethacin were replaced. The cells grown in normal media were considered control. The cell differentiation was investigated three weeks after the beginning of the culture, using oil red and alizarin red stains.

Tab. 1. Primers specifications.

| Primer | Sequence | Size (bp) | TM |
|--------|----------|----------|----|
| SPP1   | F: 5’-GGGAGTTCTCCTCGCTGATAT-3’ | 150bp | 59.35 |
| R: 5’-AGTTTGTTGCTGAATGCGCC-3’ | 57.30 |
| ALP1   | F: 5’-GGACCTTGGCTTACCAACCTC-3’ | 245bp | 61.40 |
| R: 5’-AACCTTGTCATCTCCAGGGC-3’ | 59.35 |
| BGLAP  | F: 5’-GAAATAGACTCCCGCGCTACC-3’ | 229bp | 61.40 |
| R: 5’-TCGAGTCTCGAGAGTGC-3’ | 61.40 |
Fig. 1. Preparation and surgery of rats.

Fig. 2. The steps of femur and tibia extraction from rat, primary culture of mesenchymal cells, and subculture of mesenchymal cells in 25- and 75-ml culture flasks (100, 200 and 400 \( \times \)) (Scale bar: 1 μm).
PCR confirmation of differentiation of bone marrow mesenchymal cells into osteocytes

The primers were designed using NCBI and Primer 3 software and synthesized by Sinaclon Co. (Iran) in lyophilized form. During PCR reaction (19, 20), primers are bonded on both sides of the target sequence and allow the polymerase activity of DNA polymerase. Therefore, the most important criterion for selecting a primer is that of attention to the matching of sequence of primer with template. Having the target sequence makes it possible to determine the primers sequences. The specifications of the synthesized primers are shown in Table 1.
Preparation and surgery

All experiments were carried out according to the instructions of the Animal Ethics Committee of Baqiyatallah University of Medical Sciences. This protocol relates to the observance of the ethical principles and protection of animals used for laboratory and other scientific purposes. Type I collagen was extracted from the rat’s tail. To prepare the hydrogel, 0.04 mg/ml of type 1 collagen was dissolved in acetic acid, and then mixed with cell culture media (DMEM 1X, 350 μm), FBS (10 %) and NaOH 0.4 N (26 μm). NaOH neutralizes the pH of scaffold close to that of the body. Then it was mixed with 100,000 extracted cells associated with each group and injected into the damaged area. The present study was conducted on 24 adult male rats weighing approximately 150–200 g. The rats were kept under the same conditions of 12 hours of light and 12 hours of darkness at 20–25 °C. Animal modeling was conducted by adding 100 μl nicotine in 10 ml of saline and then 200 μl was used for each rat (three times every two days). The animals had free access to water and food. Rats were anesthetized by ketamine (90 mg/kg) and xylazine (10 mg/kg) intraperitoneally. After shaving the thighs, the area was cleaned with iodine and alcohol. The hands and feet of the animals were fixed on the specified areas on the surgery bed, and then a cut was created on the skin. The fascia and muscles were cut slowly and the relevant area appeared. Then, a lesion was created using a trephine (4 mm

Fig. 4. BMSCs differentiation to osteocytes and adipocytes. Alizarin red and oil red staining were used for osteogenesis and adipogenesis differentiation confirmation, respectively.

Fig. 5. Expression of alkaline phosphatase (A), osteocalcin (B), osteopontin (C) genes in the differentiation of bone marrow mesenchymal cells. L: ladder, -: negative control, +: positive control, S: bone marrow mesenchymal cells.
in diameter), and the cells in each group were cemented to the critical size for bone regeneration. The differentiated osteocytes from bone marrow mesenchymal cells were injected to the control group (without nicotine) and those of nicotine model. The skins were stitched up by suture (4-0), disinfected with tetracycline spray and recovered. Then amoxicillin (0.1 mg/kg) was injected. After surgery, the animals were kept in a clean cage and transferred to the animal house (Fig. 1). The rats were kept inside the cage at a suitable temperature and appropriate nutrition conditions. After 4 weeks, the rats were sacrificed with a high dose of anesthesia and the specimens were removed.

**Radiographic examination**

Four weeks after the cell transplantation, radiographic examination of the right femoral bone was performed with a Senographe 600T Senix H.F device at a dose of 22 KV and 9 mas. The test was performed from the anterior-posterior (AP) and lateral (Lat) views and the rate of bone formation was scored by three persons upon inspecting the images.

**Hematoxylin-eosin staining**

For histopathologic examination, the bones were immersed in 10 % nitric acid for 2 weeks and tested with syringe tip. At the end of specified time, the samples were sent to the histopathologic laboratory, where decalcification, tissue-processing and sectioning steps (5 microns) were performed. The specimens were stained with hematoxylin-eosin (H&E) and examined and photographed with a light microscope (21–23).

**Results**

**Isolation and culture of bone mesenchymal cells**

As stated in the introduction, the characteristics of mesenchymal cells such as easy isolation and proliferation when compared to other types of stem cells, and their potential for conversion
into different types of cells led to the selection of these cells for culture and evaluation of the damaged tissue repair. For this purpose, the bone marrow mesenchymal cells isolated from rat femur and tibia were cleaned from other impurities through several steps of centrifugation with PBS and filtration. For separation of polymorphonuclears, they were added into the tube, centrifuged and then cultured. Since the mesenchymal cells typically bind to the surface, the other cells were washed out by intermittent media changes. After 4 subcultures, the cells were relatively purified and a part of them was stored and frozen (Fig. 2). Flow cytometry graphs showed that the cells with very low expression of CD34 and CD45 and high expression of CD90 and CD106 markers were mesenchymal stem cell types (Fig. 3). The mesenchymal stem cells differentiated into osteocytes and adipocytes were recognized by alizarin red and oil red, respectively. Their differentiation is shown in Figure 4.

**PCR results**

The results of PCR confirmed the expression of specific osteocyte genes including those for alkaline phosphatase, osteocalcin, and osteopontin (Fig. 5) from bone marrow mesenchymal cells differentiated into osteocytes.

**Macroscopic radiographic images**

The results of radiography showed that the rate of osteogenesis in the non-nicotine group was much higher than that in the nicotine group and the level of this density was higher in the group with differentiated bone marrow. This issue was completely true for the nicotine group. The self-repair was obviously evident in the control group without any cell transplantation compared to the nicotine group (Fig. 6).

**Microscopic histopathologic images**

The results of histologic staining showed that the rate of osteogenesis and bone formation in the canal was more pronounced in the non-nicotine group than in the nicotine group. As shown in Figure 7, the level of regular bone formation in the group differentiated with BMSCs was higher and more intensive than in other groups. On the other hand, in the control group, the process of osteogenesis toward the center and observed in the margin of fracture was also more intensive than in the nicotine control group (Fig. 7).

When comparing nicotine and non-nicotine groups, the regular and continuous bone formation in the non-nicotine group differentiated with BMSCs was more distinct than in the nicotine group in the same situation. The microscopic observations shown in Figure 8 are based on the average repair percentage in the fractured area (samples with severe fracture were not reported and not entered in the percentage determination).

**Discussion**

The bone injury is one of the most common causes of morbidity and disability in elderly patients and leads to a decline in general health and quality of life. It has been estimated that every year two million people in the United States suffer from bone fractures due to osteoporosis (24, 25). Today, restorative medicine is divided into two parts, namely cell therapy and tissue engineering. Tissue engineering is one of the subjects that can solve the crisis of tissue fracture or the problem with the shortage of organs for transplant. The cells, scaffolds and growth factors are the three major components of tissue engineering. The cells must synthesize the background material of the new tissue, the scaffold provides a suitable environment for the survival and more efficient cell function, and growth factors facilitate cellular progression to revive the new tissue. Despite several studies conducted in order to reconstruct various tissues, there are still outstanding issues such as selection of cellular sources, construction of scaffolds, cell culture, culture media, mechanical properties of cell-scaffold and appropriate animal models in this process. Therefore, the major challenges are those pertinent to cell technology, technology of structure, and attachment of designed structures in the living system (4, 5, 26, 27).

In this research, mesenchymal stem cells were isolated from the rat bone and identified by different tests such as adhesiveness, flow cytometry, differentiation of adipocytes and osteocytes by specific stains, and expression of the desired genes by PCR. The fundamentality of cells was proved and the cells were then stored for the next steps. The cells extracted from each group were differentiated into osteocytes (the third subculture cells, which were placed in osteogenesis differentiation media for 3 weeks underwent morphological changes. They got denser in some places. As a result of staining, the extracellular material in the osteocyte differentiation culture turned red). The cells were then assembled with the prepared collagen to be transferred into modeled and non-modeled animals. In this regard, in a study by Nather and colleagues, autologous mesenchymal stem cell transplantation was used to repair the femoral bone fracture in rabbits. As a result of following allograft transplantation in this study, the bone repair improved and there was an increase in osteocytes in the diaphyseal region (28). The study of Kon and colleagues was focused on bones of large animal models. They evaluated and compared healthy bones with
Effects of Bone Marrow

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