**Ex vivo Expansion and Differentiation of Mesenchymal Stem Cells from Goat Bone Marrow**

*1Mohamadreza Baghaban Eslaminejad, 1Hamid Nazarian, 1Fahimeh Falahi, 1Leila Taghiyar, 2Mohamad Taghi Daneshzadeh

**Abstract**

**Objective(s)**
Mesenchymal stem cells (MSCs) from large animals as goat which is genetically more closely related to human have rarely been gained attentions. The present study tried to isolate and characterize MSCs from goat bone marrow.

**Materials and Methods**
Fibroblastic cells appeared in goat marrow cell culture were expanded through several subcultures. Passaged-3 cells were then differentiated among the osteogenic, adipogenic and chondrogenic cell lineages to determine their MSC nature. Differentiations were determined by RT-PCR analysis of related gene expression. To identify the best culture conditions for propagation, passage-3 cells were plated either at varying cell densities or different fetal bovine serum (FBS) concentrations for a week, at the end of which the cultures were statistically compared with respect to the cell proliferation. In this study, we also determined goat MSC population doubling time (PDT) as the index of their in vitro expansion rate.

**Results**
Passage-3 fibroblastic cells tended to differentiate into skeletal cell lineages. This was evident in both specific staining as well as the specific gene expression profile. Moreover, there appeared to be more expansion when the cultures were initiated at 100 cells/cm² in a medium supplemented with 15% FBS. A relatively short PDT (24.94±2.67 hr) was a reflection of the goat MSC rapid rate of expansion.

**Conclusion**
Taken together, fibroblastic cells developed at goat marrow cell culture are able to differentiate into skeletal cell lineages. They undergo extensive proliferation when being plated at low cell density in 15% FBS concentration.

**Keywords:** Adipogenesis, Bovine serum, Cell seeding density, Chondrogenesis, Goat mesenchymal stem cells, Osteogenesis

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1- Stem Cell Department, Cell Science Research Centre, Royan Institute, ACECR, Tehran, Iran

*Corresponding Author: Tell: +98-21-22413790; Fax: +98-21-22409314; email: eslami@royaninstitute.org

2-Royan Animal Facility, Karaj, Iran
Introduction
Mesenchymal stem cells (MSCs) possess two fundamental characteristics: the ability of extensive replication and the capacity of multilineage differentiation among bone, cartilage and adipose cell lineages. These cells have been reported to be present in bone marrow and several other tissues in adult (1). MSCs were first described by Friedenstein and co-workers who have reported the isolation of the cells from rat bone marrow tissue, using the cell plastic-adherent property (2). Since then, this property has been used to isolate and characterize MSCs from a number of species including dog, rabbit, pig, baboon and human (3-7). Having the ability of self-renewal for long-term along with multilineage differentiation potential have rendered MSCs as significant cellular materials for tissue regeneration applications (8).

Two important challenges associated with MSCs would be the topics of their safety and efficacy in use as therapeutic tool for tissue regeneration. These subjects need sufficient investigation in animal models, which in turn requires MSCs isolation and characterization, as the first step, from the interest animal. In this regards, most related investigation has so far been carried out in small animal as rodents (9). MSCs from large animals as goat which is more similar to human than rodents has rarely gained attention. MSCs biology in this animal has indeed remained to be fully investigated. In those research works associated with goat MSCs, the cells isolated from the animal marrow have mostly been used to establish 3D cultures in scaffolds without taking into account the cell in vitro characteristics as well as culture requirements. In this regards, Nair et al (2008) have manufactured a triphasic scaffolds consisting of hydroxyapatite, tricalcium phosphate and calcium silicate and evaluated the scaffold biocompatibility with goat MSCs 3D culture (10). In a similar investigation, Li and colleagues have used goat MSCs for biocompatibility test of newly manufactured hydrogel scaffolds (11). Dai et al have investigated the chondrogenic potential of goat MSCs within hydrogel in vitro (9). Williams et al have used goats as animal model to evaluate the regenerative capacity of goat MSCs-loaded scaffolds in bone large defects (12). Neither of these experiments has examined the multipotent differentiation capacity of the isolated cells which MSC committee of international society of cell therapy has announced it as defining criterion for MSCs (13). In the present study, this subject was examined.

Carrying out the differentiation analysis with RT-PCR, we noticed that a number of specific genes for goat bone, cartilage and adipose cell were not registered in gene database of NCBI. As a solution, we designed and used the primers for equine corresponding genes (which were available in gene database) to amplify the goat genes; and then the amplification product was extracted, sequenced and investigated whether or not they were similar to corresponding genes in close species to goat. All sequences were then registered in NCBI gene database. Accession numbers were assigned to the sequences which are currently available. Moreover, in this study, culture conditions for the maximum proliferation of goat MSCs were optimized.

Materials and Methods
Bone marrow cell culture
Four male adult goats of Najdi strain were used in this study. Before experiments, approval of the animal use was obtained from the Ethic Committee of Royan Institute. The animals were kept in the facility of Royan Institute (Karaj, Iran) till become acclimatized to housing and diet. Under general anesthesia by Ketamine (22 mg/kg) and xylazine (0.2 mg/kg) IM injection, bone marrow aspirate (about 10 ml) were drawn from the tibia, collected into 50 ml tube containing 7500 unit heparin and shipped on the ice to cell culture facility of Royan Institute. Bone marrow aspirates were added to 5 ml DMEM medium (Dulbecco’s modified eagle medium, Sigma, USA) supplemented with 100 IU/ml penicillin (Sigma, USA), 100 IU/ml streptomycin...
(Sigma, USA), 10% FBS (Fetal Bovine Serum) and centrifuged at 300 g for 5 min. Supernatant was discarded, cell pellet resuspended in 1 ml DMEM medium and plated at $5\times10^6$ cells/cm$^2$ in 75-cm$^2$ culture flasks in an atmosphere of 5% CO$_2$ and 37 °C. About 3-4 days after culture initiation, culture medium was replaced with fresh DMEM and the flasks were incubated with a medium replacement of 2 times a week until the cultures became confluent. At this stage, cells were trypsinized and subcultured at $5\times10^5$ cells/cm$^2$. Cell passages were repeated up to the subculture 3 when sufficient cells were available to continue the experiments described below.

**Differentiation potential**

To evaluate the mesenchymal stem cell nature, the isolated cells were differentiated into osteogenic, chondrogenic and adipogenic cell lineages.

To induce osteogenic differentiation, confluenced passaged-3 cells were cultured in the DMEM medium supplemented with 50 mg/ml ascorbic 2-phosphate (Sigma, USA), 10 nM dexamethasone (Sigma, USA) and 10 mM β-glycerol phosphate (Sigma, USA) for 3 weeks (14-15). At the end of this period, alizarin red staining was used to observe the matrix mineralization. For staining, the cultures were first fixed by methanol for 10 min and then subjected to alizarine red solution for 2 min.

For adipogenesis, DMEM medium containing 100 nM dexamethasone (Sigma, USA) and 50 mg/ml indomethacin (Sigma, USA) was used to induce the differentiation in the confluent culture of passaged-3 cells (14, 15). Three weeks after culture initiation, the cells were fixed with 4% formalin at room temperature, washed by 70% ethanol and stained, using oil red solution in 99% isopropanol for 15 min.

To induce the cartilage differentiation, micro mass culture system was used. For this purpose, $2.5\times10^5$ passaged-3 cells were pelleted under 300 g for 5 min and cultured in a DMEM medium supplemented by 10 ng/ml TGF-ß3 (transforming growth factor-ß3) (Sigma, Germany), 10 ng/ml BMP6 (bone morphogenetic protein-6) (Sigma, Germany), 50 mg/ml insulin transferin selenium+ premix (Sigma, Germany) and 1.25 mg bovine serum albumin (Sigma, Germany) and 1% fetal bovine serum (15, 16). Three weeks after initiation of the culture, the pellets were removed and subjected to the following: fixing in 10% formalin; dehydrating in ascending concentrations of ethanol; clearing in xylene; embedding in paraffin wax and sectioning at 5 µm by microtome. The sections were then stained, using toluidine blue for 30 sec at room temperature.

**RT-PCR analysis**

**Primers**

Because of the lack of a number of specific gene sequences for goat osteocyte, chondrocyte and adipocyte (except for GAPDH and Osteocalcin), we designed the primers for equine bone, cartilage and adipose-specific genes which were available in NCBI gene Database and used them in order to amplify the genes in goat differentiated cultures. The bands of amplified fragments were recovered from the gel by QIA quick Gel Extraction kit and then sequenced by Chromas Software version 2.31. BLAST results of these sequences, showed that there are many similarities between them and sequences related to the genes of other species close to goat. Therefore, it confirmed that the equine specific primers were able to amplify our interest fragments in goat genome. Sequenced gene fragments were registered in NCBI gene database and they were assigned the accession numbers. The similarity percentage of these sequences to the sequences of other species and also primer sequences used to amplify the interest genes are indicated in Table 1 and 2, respectively. The sequences that have been registered at NCBI gene database for goat are shown in Table 3.

| Table 1. The similarity percentage of amplified sequences from ovine differentiated cultures, using equine primers compared with those of other species. |
Goat Mesenchymal Stem Cell

<table>
<thead>
<tr>
<th>Gene name</th>
<th>Other species</th>
<th>Similarity percentage</th>
<th>Other species</th>
<th>Similarity percentage</th>
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<td>Equus caballus</td>
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<tr>
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<tr>
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Table 2. Primers used in RT-PCR analysis, * indicates the genes that were not found in NCBI gene database. These genes were sequenced in this study and registered at NCBI gene database. Their accession numbers indicated in the left column are now available in the database.

<table>
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<td></td>
<td></td>
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RT-PCR Procedure

Total RNA was isolated, using the RNX™ (-Plus) (RN7713C; CinnaGen Inc., Tehran, Iran). Before RT, a sample of the isolated RNA was treated with 1 U/ml of RNase-free DNase I (EN0521; Fermentas, Opelstrasse 9, Germany) per 1 mg of RNA in order to eliminate residual DNA in the presence of 40 U/ml of ribonuclease inhibitor (E00311; Fermentas, Germany) and 1×reaction buffer with MgCl₂ for 30 min at 37 °C. To inactivate the DNase I, 1 ml of 25 mM EDTA was added and incubated at 65 °C for 10 min. Standard RT Reactions were performed with 2 μg total RNA, using Random hexamer as a primer designed for equine genes and a RevertAid™ First Strand cDNA Synthesis Kit (K1622; Fermentas, Germany) according to the manufacturer’s instructions. For every reaction set, one RNA sample was prepared without RevertAid™ M-MuLV Reverse Transcriptase (RT - reaction) to provide a negative control in the subsequent PCR. To minimize variation in the RT reaction, all RNA samples from a single experimental set up were reverse transcribed simultaneously. Reaction mixtures for PCR included 2 ml cDNA, 1×PCR buffer (AMS™, CinnaGen Co., Tehran, Iran), 200 mM dNTPs, 0.5 mM of each antisense and sense primer, and 1 U Taq DNA polymerase. PCR was performed, using Mastercycler Gradient 5331, Ependorf, Hamburg, Germany.

Table 3. The sequences of goat bone, cartilage and adipose genes that have been registered in NCBI gene database by the authors. The Accession numbers that have been devoted to these sequences are indicated in Table 2.
Optimizing the culture condition for maximum proliferation

In this study, initiating cell seeding density and FBS concentration in medium were optimized for extensive proliferation of the cells. For this purpose, the isolated cells (passage-3) were plated in a 100-mm dish at varying densities of 10, 50, 100, 200 and 500 cell/cm² in DMEM, supplemented with 100 IU/ml penicillin, 100 mg/ml streptomycin and 10% fetal calf serum for a week. At the end of this period, the cells were lifted, counted and the fold increase number was calculated and compared statistically (One-way ANOVA).

Likewise, the cultures of passage-3 cells were established with 100 cell/cm² in DMEM, supplemented with such varying concentrations of FCS as 5%, 10%, 15% and 20% for one week. Subsequently, the fold increase number of the cells was calculated in each group and compared by means of One-way ANOVA.

CFU-F (colony forming unit-fibroblast) assay

Hundred cells of the passage-3 culture were plated in a 25-cm² plastic flask in DMEM, supplemented with 100 IU/ml penicillin, 100 mg/ml streptomycin and 10% fetal calf serum before they were incubated at 37 °C and 5% CO₂ for 6 days. At the end of the cultivation period the number of the clones was counted under an inverted microscope.

Population doubling time (PDT)

After the optimal cell seeding density and FBS concentrations were determined, the cells cultivated under these conditions and their PDT was calculated. For this purpose, passaged-3 cells were plated at 100 cells/cm² in a DMEM supplemented with 15% FBS and incubated for a period of 7 days, when the cultures terminated, the cells lifted and counted with hemocytometer. Using the following equation, PDT was determined. In this equation N is the initiating cell number, N₀ the harvesting cell number and CT stands for culture time.

\[
(PDT) = \frac{CT}{\log \frac{N}{N_0} \times 3.31}
\]

Results

Cell culture

The primary cultures of the goat bone marrow cells contained mainly fibroblast cells as well as a few small round cells. The number of latter was reduced by performing subcultures, during which the fibroblastic MSCs were purified and became dominant in the culture (Figure 1).
Goat Mesenchymal Stem Cell

Figure 1. Goat marrow cell culture. At primary culture, fibroblastic as well as small clear cells can be observed. The number of clear cells was decreased during the passages and fibroblastic cells became dominant in culture.

Multi-lineage differentiation
In osteoinductive cultures, in some areas, nodule-like structures were observed. Following alizarin red staining, red mineralizing areas of cultures appeared (Figure 2A). Differentiation was further demonstrated by RT-PCR analysis of the osteocytic markers. After a 3-week induction period, osteocalcin and collagen I and III mRNA levels were largely produced (Figure 2B).

Nodular structures were not observed at adipogenic culture. Small lipid droplets appeared within the cytoplasm of the cells first on days 3 to 4; they gradually occupied the whole cells by day 21. The lipid droplets turned into red when stained by the oil red staining method (Figure 2C). RT-PCR analysis was also indicative of the expression of PPAR alpha (Peroxisome proliferator’s activated-receptor alpha), PPAR gamma (Peroxisome proliferator’s activated-receptor gamma), LPL and LPL (lipoprotein lipase) genes (Figure 2D).

In the micro mass culture system for chondrocyte differentiation, the size of the pellet seemed to have increased during the culture period, probably as a result of matrix production and secretion. Metachromatic nature of the matrix was demonstrated by the toluidine blue staining method (Figure 2E), and its marker molecules including aggrecan, Collagen II and Collagen X were examined by RT-PCR analysis (Figure 2F).

Figure 2. Multilineage differentiation potential of the isolated cells. Passage-3 cultures were easily differentiated among bone (A-B), cartilage (C-D) and adipose (E-F) cells. Differentiation into the skeletal cell lineage is considered to be as defining criterion for the cells.

Culture optimization
Seeding density
The maximum fold increase in cell number was achieved when the cells were plated at 100 cell/cm² (Figure 3, upper graph). At this point, their difference in comparison to the groups of 10, 50, 200 and 500 cell/cm² was statistically significant ($P<0.01$).
Figure 3. Optimization of the culture conditions for extensive proliferation of the cells. Passage-3 goat MSCs possessed maximum proliferation when they were plated at 100 cells/cm² (upper graph) in a medium supplemented with 15% FBS (lower graph). * indicates a statistically higher value (P<0.05).

Serum concentration
The cells having been plated with 15% FBS had a significantly larger fold increase in cell number as compared to those having been plated with 5%, 10% and 20% FBS (Figure 3, lower graph).

CFU-F Assay Results
Almost 58.8 clones, each consisting of 20-70 cells, formed by per 100 cells plated in 25-cm² flasks for 6 days.

Population doubling time (PDT)
Population of passaged-3 goat MSCs appeared to have a rather high expansion rate, so that their number was doubled in average of 24.94±2.67 hr when plated at 100 cells/cm² in a medium supplemented with 15% FBS.

Discussion
In this study, we culture-expanded the cells from goat marrow and evaluated them whether or not they were able to produce specialized lineages as bone, cartilage and adipose cells. The studied cells appeared to be mesenchymal stem cells in nature because they successfully differentiated into the skeletal cell lineages. It should be mentioned that since investigators have reported studies of MSC, using different methods of isolation and expansion, and different approaches to characterizing the cells, thus it is increasingly difficult to compare and contrast study outcomes which hinders progress in the field. To address this issue mesenchymal and tissue stem cell committee of international society for cellular therapy has proposed the ability of multilineage differentiation as defining criterion for MSCs (13). In a few related investigations that have previously been conducted, these important defining characteristics have been ignored; hence, it was not clear that the cells having been used by those researchers were the MSC in nature. In the present study multilineage differentiation capacity of marrow-derived goat MSCs were examined and confirmed.

In addition to its significance as a defining criterion for MSCs, in vitro differentiation of MSCs would be of great clinical value. In cell therapy strategies; it is believed that transplanted cells should be of terminally differentiated lineages otherwise they would produce non specific cells in defect site (17-18). Therefore, in vitro differentiations of the cells into interest cells could be a necessary step prior to their transplantation. In this study the potential of goat MSC in vitro differentiation which has not gained attention at former investigations were explored.

The other point of the present study was the issue of the lack of a number of goat bones, cartilage and adipose cell-specific gene sequences in NCBI gene database. To solve this problem, we used the primers designed for equine corresponding genes and amplified the genes in differentiated cultures of goat MSC and then extracted the products and sequenced them. Our further examination by BLAST software indicated the similarity of the sequenced products with known sequence of the genes in several species. Therefore, we concluded that they would be the sequence that would be conserved in the species. The sequences were then registered at NCBI gene
database and were gained an accession number indicated in Table 1.

MSCs are being considered as valuable tool for regenerative medicine. The preclinical development of therapeutic procedures, using MSCs requires sufficient animal studies both for safety and for the investigation of efficacy of the cells. In this regard, the investigation of MSC in large animals would be of great importance since in contrast to rodents they are genetically more closely related to humans. Results from large animal models have been much more predictive with regard to human clinical trials than have results from in vitro assays or rodent models. Furthermore, one limitation associated with mice as a model is their limited life expectancies that preclude long-term follow-up (19, 20). Given the significance of large animal model in stem cell investigation, in this study MSCs from goat bone marrow were isolated and characterized. This has not been performed at related previous investigations.

MSCs occur at low frequencies in bone marrow samples (21). On the other hand they are needed in large numbers in most experimental set up. As a solution, MSCs should be expanded in culture before they are being used at any application. MSC ex vivo propagation is largely dependent on the presence of FBS in culture medium. According to former researches, MSCs that are developed in mediums containing FBS would be immunogenic and may transfer bovine pathogens upon transplantation (19). In spite of extensive attempt that has been made to substitute the bovine serum, less success has been achieved (21, 22). In this study, therefore, goat MSCs were cultivated in different concentrations of FBS to determine the least concentration with the highest mitogenic effect, which this was achieved by 15% FBS in the medium.

In the present study, the best cell density at culture initiation which resulted in extensive proliferation of the cells was determined. This is of utmost importance especially at cell therapy strategies where the rapid expansion of cells would be desired. According to our data, goat MSCs possessed the highest fold increase when being cultivated at 100 cells/cm². In a study by Bartmann et al it has been reported that there would be an inverse correlation between human MSC seeding density and their proliferation (23). Our results seem to be differed from those of Bartmann et al in that, although, the culture with the least cell concentration showed the highest proliferations, but in culture with 2000 cells/cm² there appeared to be more proliferation than in culture with 500 and 1000 cells/cm². This discrepancy would be the results of different species from which the cells obtained (human versus goat). Such results, however, can be used to optimize the goat culture condition for maximum proliferation of the cells.

In spite of the considerable attempts that have been made to define the antigenic profile of human MSCs, no definitive single marker has so far been introduced. In this regard, several markers including CD133, LNGFR (low affinity nerve growth factor receptor) and STRO-1 have been proposed as markers of human MSCs (24-26). However, the identification of MSCs, especially those from animal source would be a difficult task owing to the lack of distinct specific marker. To overcome this problem, MSCs committee of the international society for cell therapy has proposed that MSCs from animal source should possess two characteristic criteria: a) they must be adherent on plastic surfaces of culture dish. b) They must be able to produce bone, cartilage and adipose cell lineages (13). Since the isolated cells of the present study were easily differentiated into bone, cartilage and adipose cells and also they were plastic adherent cells, we convinced that they were the MSCs described elsewhere.

Conclusion

Taken together, fibroblastic cells developed at marrow cell culture of goat are able to differentiate into skeletal linage as bone cartilage and adipose cells, hence, they are MSCs described elsewhere. According to our findings, the best culture condition for maximum proliferation of goat MSCs would
be achieved when the cells were plated at 100 cells/cm² in a medium containing 15% FBS. Furthermore, in this study, we identified the sequences of a number of goat genes which were not present in NCBI gene database. The genes were registered and gained an accession number.

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References

