Description of the technique of bone marrow harvesting in the coxal tuberosity for isolation and culture of mesenchymal stem cells of buffaloes (*Bubalus bubalis*)

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**ABSTRACT**

Several studies with mesenchymal stem cells (MSCs) have been developed in many species because of its ability to differentiate into other mesoderm lineages, capacity of self-regeneration, low immunogenicity, paracrine, anti-inflammatory, immunomodulatory and antiapoptotic effects which make them a promissory source to be used in therapeutic strategies. The aim of this study is to report the technique of harvest of bone marrow (BM) in the coxal tuberosity (CT) of buffaloes and its processing and cultivation. For this, after anesthetic block from the region corresponding to the CT, bone marrow harvesting was performed with a myelogram’s needle. The samples collected showed plastic adherence with 96 hours and took approximately 32 days to reach 80% confluence. And then differentiation into adipogenic and osteogenic lineages was performed. Samples showed morphological changes during differentiation protocol, but not all presented production of extracellular deposits of calcium or intracellular fat droplets. The anatomical site tested showed to be an alternative site of harvest of BM once provided with the appropriate isolation and culture of the mononuclear fraction. Moreover, the procedure was performed without difficulty and with great security. Based on this, it can be concluded that CT is an excellent anatomical site for isolation and culture of MSCs and the proposed technique is viable and feasible to be held in buffaloes.

**KEYWORDS**

Buffaloes; MSCs; Coxal Tuberosity; Culture

1. INTRODUCTION

The ability of mesenchymal stem cells (MSCs) to differentiate into other mesoderm lineages such as bone and cartilage opened a variety of experimental strategies to investigate the possibility of these cells to be used in tissue engineering since MSCs derived from bone marrow (BM) have been applied in the treatment of musculoskeletal disease in several species [1]. Besides the differentiation capacity, those cells have the capacity of self-regeneration, low immunogenicity, paracrine, anti-inflammatory, immunomodulatory and antiapoptotic effects which make these cells a great alternative to be used in regenerative therapies.

Bone marrow-derived mesenchymal stem cells (BM-MSCs) are the most studied source of MSCs and because of this, they have received special attention and are best characterized [2]. Bone marrow harvesting on coxal tuberosity (CT) has already been described in other species such as horses [3,4], dog, cat [5], sheep [6], pig [7] and humans [8].

In buffaloes, there are reports of studies with embryonic stem cells [9], MSCs derived from adipose tissue [1] and from amniotic fluid [10]. Based on this, this work...
aims to evaluate an alternative site of harvest of bone marrow in the CT and the cultivation of mesenchymal stem cells. The results of this study can contribute to and aid in the harvest of buffalo MSCs and maximize their therapeutic use.

2. MATERIALS AND METHODS

2.1. Animals

Six healthy adult female buffaloes aged 8 - 14 years were selected and used. The experimental protocol (No. 160/2012- CEUA) was approved by the ethics and welfare committee of the Sao Paulo State University - Botucatu. All procedures were performed under the international guidelines for the care and use of experimental animals.

2.2. BM Harvesting

Buffalo BM harvesting was performed on selected animals properly contained in a stock. The anatomical site of right CT was properly identified by palpation and it was performed trichotomy of the region. After this, antisepsis was done with chlorhexidine and alcohol (Riohex, Rioquimica—Brazil) followed by local anesthetic block with 10 ml of 2% lidocaine (Cristalina, Brazil). Past ten minutes, a myelogram’s needle (Lang®—Brasil) was introduced approximately 2 cm for bone marrow harvesting at an angle of approximately 45 degrees relative to the animal in dorsoventral and cranio-caudal direction (Figure 1). Once the needle was firmly fixed in the CT, the mandril was removed and proceeded BM aspiration (Figure 2) with a syringe of 20 ml properly identified containing 1 ml of heparin at 1000 IU/ml (Hemofol®, Cristalina, BRA) and 1 ml of PBS phosphate-saline, pH = 7.2 (PBS® 1x, LGC Biotechnology, BRA). After BM harvesting, each sample collected was manually homogenized, identified, cooled and referred to the laboratory for isolation, cultivation and characterization from mononuclear fraction obtained from Histopaque® (1077—Sigma®, USA) gradient in a laminar flux environment.

2.3. Isolation, Culture and Differentiation of MSC

After collection, the samples were filtered and centrifuged at 340 g for ten minutes. The supernatant was removed and culture medium composed by DMEM low-glucose (Invitrogen®, USA) was added.

Mononuclear fraction (MF) was isolated in a Histopaque® (1077—Sigma®, USA) gradient (Figure 3), trough 40 minutes centrifugation at 340 g. The MF was washed twice and cultured in DMEM low glucose/F12 (1:1), supplemented with 10% fetal bovine serum,
antibiotics (1%) and antimycotics (1.2%) (Invitrogen®, USA) in humidified atmosphere containing 5% CO₂. Maintenance medium was changed at each two or three days and when the cells reached 80% confluence, first passage was performed by resuspending the cells with TrypLE Express® (Invitrogen®, USA, cat. number: 12604021), and transfer to 6-wells plates (Sarstedt®, USA) for differentiation assay into osteogenic and adipogenic lineage.

The differentiation protocol took about 10 days using media and supplements prepared from the Adipogenesis Differentiation Kit (Invitrogen®, USA, cat. number: A10070-01) and the STEMPRO® Chondrogenesis media and supplements prepared from the Adipogenesis.

Differentiation Kit (Invitrogen®, USA, cat. number: A10070-01) and the STEMPRO® Chondrogenesis Differentiation Kit (Invitrogen®, USA, cat. number: A10071-01) according to manufacturer’s recommendation. The medium of differentiation was changed every two days. At the end of differentiation period the morphological changes of cells were analyzed and samples were stained with 2% Alizarin Red (Sigma®, USA, pH = 4.2) and 0.5% Oil Red (Sigma®, US), for the observation of the deposit of calcium in the extracellular matrix and the presence of intracytoplasmic lipids droplets, respectively.

3. RESULTS

BM harvesting on CT was easily conducted and it was not found any difficult or problem during or after the collection. Furthermore, the needle, anesthetic protocol, doses and MSCs processing was considerate appropriate for the species under study.

From each animal it was collected on average 3 ml of BM and this volume was sufficient for the processing of the sample from these animals, promoting correct isolation, culture and characterization of the cells. In one animal, with advanced age, the volume collected was lower (1 ml) and the culture failed. The process of cooling the material during transport apparently did not affected the quality of the samples once all samples were submitted to culture.

The adhesion time for these cells was four days and the period between the adhesion and 80% confluence was on average 32 days (Figure 4). Morphological changes in cells in culture were observed during the differentiations period (Figure 5). Of all samples collected, the culture of only one animal (the older) failed. So of the five samples, four (80%) were able to differentiate in osteogenic lineage and two (40%) were able to differentiate in the adipogenic lineage. All samples showed morphological changes, but some presented no production of extracellular deposits of calcium or intracellular fat droplets, observed by staining with Alizarin Red and Oil Red, respectively.

4. DISCUSSION

The collection of BM in the CT of buffalo is considered safe both for the veterinarian and for the animal. Once held the correct contention, fact was also reported in horses [3].

Despite the possibility of obtaining MSCs from other sources such as adipose tissue [1], umbilical cord blood [8], peripheral blood [11], amniotic fluid [10], BM is considered the one that has the most abundant source of MSCs [8].
For BM harvesting in the CT of buffaloes, there is no need for general anesthesia, once a simple anesthetic block already allows the development of the technique. Such a fact makes the technique advantageous since adverse effects have been reported in human patients undergoing general anesthesia such as pain at the site of collection or even anemia [11]. Besides, this harvest can be done with the live animal, so there is no need to make a surgical wound. And the cells can be used in the same animal.

In this experiment, the volume collected from all animals was considerate properly. However, in other species, the volume collected was greater with data of 60 ml in pigs [12], 10 ml in humans [8], 11 ml in horses [3] or 1 ml in dogs [13]. As already reported, the best sample for isolation and expansion of MSCs is obtained in the early stages of bone marrow aspiration. Furthermore, it is known that the volume of material obtained decreases along the samples [8].

In one animal from this study, it was observed that the volume material collected was lower (1 ml) and yellowish. This can be explained by the advanced age of this animal in order that the hematopoietic tissue is gradually replaced by non-hematopoietic mesenchymal cells, also called fatty bone marrow [14].

Compared with samples from other species processed in the same laboratory, a longer primary culture time was observed during culture. For example, in dogs the time to reach confluence is on average 10 days (unpublished data) and in horses is on average 22 days [15].

With respect to differentiation, heterogeneity in response to differentiation after induction was noted. Perhaps this response is linked with an inadequacy of the differentiation media used. However, it is important to remember that intrinsic difference among the samples cannot be ruled out. Since the population of mesenchymal stem cells contained in bone marrow environment is heterogeneous, it is committed to diverse differentiation processes.

With this protocol proposed, it was evident that the antisepsis, local block and site of harvest were efficient on buffaloes and can be used in this species for bone marrow harvesting from CT. The anatomical site tested showed to be an alternative site of harvest of BM once provided with appropriate isolation, culture and characterization of mononuclear fraction. Besides, the procedure was performed without difficulty and with great security. Although the osteogenic potential was evident, the adipogenic differentiation needs to be improved. Based on the above, it can be concluded that CT is an excellent anatomical site for isolation and culture of MSCs and that the proposed technique is viable and feasible to be held in buffaloes and can be used in future researches.

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REFERENCES


