INTRODUCTION

Lameness due to musculoskeletal injuries is the most frequent problem in the equine industry and leads to substantial financial losses related to treatment, resting periods, recurrence and poor performance in athlete equines (Dyson 2004). Most of these injuries are related to musculoskeletal structures, such as joints, tendons and ligaments, which generally respond to conventional therapies with scar tissue formation leading to chronic and/or degenerative processes, ending in a high risk of recurrence. The application of regenerative medicine for the treatment of musculoskeletal diseases has widely increased in veterinary medicine due to the perspective of higher successful rates (Stewart and Stewart 2011, Owens et al 2011). The most common related techniques consider the application of platelet rich plasma (PRP) and/or mesenchymal stem cells (MSCs). The main sources of MSCs in the clinical setting are: bone marrow (BM), adipose tissue and umbilical cord blood (Taylor and Clegg 2011). From BM, there are two main products that...
are being currently applied for the treatment of orthopedic injuries in the horse: bone marrow mononuclear cell fraction (BMMNCs), which comprise some hematopoietic progenitor cells, MSCs and monocytes, which can be readily isolated and injected into the lesion; on the other hand, there is the use of cultivated MSCs derived from the mononuclear cells (MNCs) fraction of BM. Both have demonstrated to enhance the repair of soft tissue injuries in horses (Smith et al 2003, Oliveira 2008, Crovace et al 2010), nonetheless, all the described protocols require access to fully equipped and costly laboratory facilities. Besides, obtaining a pure fraction of cultivated MSCs also demands longer time (2-3 weeks) (Fortier and Smith 2008, Kol et al 2013) and higher costs related to cell culture (Owens et al 2011). This study aims to demonstrate to equine and other veterinary practitioners that obtaining and isolating BMMNCs from horses is a simple procedure and can be performed with basic laboratory equipment.

MATERIAL AND METHODS

All necessary equipment should be assembled before beginning the procedure (table 1). There is a variety of needles available to aspirate BM in the horse. The authors prefer to use an 11G x 10 cm Komiyashiki reusable needle (figure 1a) for this purpose.

HORSES

In order to demonstrate the applicability of this technique, the authors performed this procedure in 10 healthy crossbred adult horses ranging between 5-17 years of age.

BONE MARROW ASPIRATION

The horses were appropriately restrained using stocks and sedation according to their temperament (Taylor and Clegg 2011). The administration of xylazine1 (0.5-1 mg/kg, IV) provides sufficient restraint for most adult horses. Its combination with an opiate (butorphanol 0.02 mg/kg IV or morphine 0.1 mg/kg IV) or the use of detomidine (0.02 mg/kg, IV) (Taylor and Clarke 2007) is recommended for patients that require stronger sedation (Kasashima et al 2011). A 10 x 10 cm wide band overlying the ventral midline in the cranial thorax at the level of the elbows and the 5th sternebra was located (Kasashima et al 2011). Local anaesthetic (10 mL, 2% mepivicaine solution) was injected at the predicted aspiration area and scrubbed a final time.

Prior to aspiration, one 60 mL syringe was aseptically preloaded with 15000 IU of heparin + 10 mL (DPBS) and preserved sterile. An 11G x 10 cm Komiyashiki needle was introduced perpendicularly through the anaesthetised area and moved forward until it contacted the middle length of the ventral midline surface of the 5th sternebra (figure 1b, 1c). The index finger was placed no more than 2 cm behind the tip of the needle, which was gradually moved forward using rotating movements until it was firmly seated into the bone to a depth of 1-2 cm into the medullar cavity (Sellon 2006, Taylor and Clegg 2011, Kasashima et al 2011). The stylet moved forward removed from the needle, the 60 mL syringe moved forward attached to its hub and gentle pulsatile aspiration was applied to obtain the sample. The sample was carefully mixed by several inversions to ensure an adequate combination of heparin and marrow that prevents clotting. One drop of the 60 mL syringe content was placed on a clear Petri dish and grossly examined for evidence of small, grayish marrow spikes and fat particles that adhere loosely to the plastic (Taylor and Clegg 2011). If spikes and fat were observed, it was likely that a sample of adequate origin had been obtained, therefore the needle could be removed and sterile gauze was held over the area with light pressure to facilitate haemostasis. The sample was transported to the on-site laboratory in cooling recipients at 5° to 8 °C (same as used for semen transportation). Generally, between 15 to 20 mL are harvested per aspirate. If the marrow sample obtained in the first attempt is not enough, needle placement should be altered to the adjacent sternebra and the procedure should be repeated.

<table>
<thead>
<tr>
<th>Table 1.</th>
<th>Equipment and supplies required to harvest bone marrow from an adult horse and isolate bone marrow mononuclear cells.</th>
<th>Equipos y materiales requeridos para obtener médula ósea de un caballo adulto y aislar células mononucleares de medula ósea external.</th>
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<tbody>
<tr>
<td>Drugs for chemical restrain</td>
<td>Clippers</td>
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<tr>
<td>Surgical prep supplies</td>
<td>Surgical prep supplies</td>
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<tr>
<td>Sterile gloves</td>
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<tr>
<td>Komiyashiki needle2</td>
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<tr>
<td>Mepivacaine3</td>
<td>Mepivacaine3</td>
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<tr>
<td>Sterile 60mL syringe containing 15000 IU sodium heparin (3 mL)</td>
<td>Sterile 60mL syringe containing 15000 IU sodium heparin (3 mL)</td>
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<tr>
<td>Plastic Petri dish</td>
<td>Plastic Petri dish</td>
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<td>Sterile 4x4 gauze pads</td>
<td>Sterile 4x4 gauze pads</td>
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<tr>
<td>Pasteur pipette / Sterile IV catheter 14Gx2”</td>
<td>Pasteur pipette / Sterile IV catheter 14Gx2”</td>
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<tr>
<td>Latex gloves</td>
<td>Latex gloves</td>
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<tr>
<td>Dulbecco’s Phosphate-Buffered Saline (DPBS)4</td>
<td>Dulbecco’s Phosphate-Buffered Saline (DPBS)4</td>
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<tr>
<td>Falcon®5 15 ml tubes</td>
<td>Falcon®5 15 ml tubes</td>
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<tr>
<td>Ficoll®6</td>
<td>Ficoll®6</td>
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<tr>
<td>Swinging bucket centrifuge</td>
<td>Swinging bucket centrifuge</td>
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</table>

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1 König Lab., Montevideo, Uruguay.
2 Fremiqsur Ltda., Temuco, Chile.
3 Pfizer, New York, USA.
4 Invitrogen Canada Inc., Burlington, Ontario, Canada.
5 BD Biosciences, Santiago, Chile.
6 Sigma-Aldrich, Santiago, Chile.
EVALUATION OF THE ORIGIN OF THE SAMPLE (OPTIONAL)

Even though objective confirmation of sample’s origin is not essential for experienced practitioners, a stained smear can be processed. For this purpose, marrow particles are collected with pipette and placed on one end of a glass slide, which is then held vertically. Particles tend to stick to the slide while blood runs off. A second glass slide was placed across the area of particle adherence, perpendicular to the first slide. After marrow spreads between the slides, they were pulled apart in the horizontal plane (Reagan et al 2011). The prepared slide was then dried quickly by waving it rapidly in the air and stained using a Romanowsky-type stain (Wright-Giemsa or Dif Quik) (Sellon 2006, Reagan et al 2011). The stained smear was scanned microscopically using a 10X and 40X objectives to ensure the presence of marrow tissue, megakaryocytes and hematopoietic precursor cells (figure 3a, 3b).

MONONUCLEAR CELLS ISOLATION

The sample should be initially filtered in a transfusion set of 200 µm in order to remove bone fragments and cell clumps (figure 2a, 2b), and then gently layered over 2.5 mL Ficoll® solution in sterile 15 mL Falcon® tube (Bourzac et al 2010). The tube was centrifuged at 500g for 30 minutes at 5˚C. After centrifugation, the supernatant plasma is carefully collected by aspiration with a sterile Pasteur pipette, and discarded, avoiding drag cells from the sediment. The clear layer containing the MNCs and Ficoll® (figure 2c, 2d), was then transferred to another sterile Falcon® tube which was completed with DPBS and centrifuged at 300 g for 10 minutes at 5˚C. Then, 80% of the supernatant was removed and the remnant completed with DPBS followed by a last centrifugation at 300 g for 10 minutes at 5 ˚C, to remove the excess of Ficoll®. Most of the supernatant (= 90%) was discarded and an aliquot of 8-10 µl of the isolated cell suspension was subjected to a cell count and viability determination.

TOTAL CELL COUNT AND VIABILITY TEST

Whereas not mandatory, in order to access the sample quality, the concentration of isolated MNCs in this study was determined using a Neubauer® chamber or a conventional blood cell automated counter. Viability can be confirmed through the application of a Dye Exclusion Test using trypan blue in a Neubauer® chamber, a simple and precise technique (Kerr 2002). A Gram staining was performed on the samples of this study in order to evaluate bacterial contamination.

FACILITIES/WORKING ENVIRONMENT

Other reported protocols for the isolation of BM products consider the use of laminar flow tissue culture hood which are sterilised with 70% ethanol or isopropanol before and after use (Helgason 2004, Taylor and Clegg 2011), however, the authors proposed that it can be replaced by good sterile technique in a clean laboratory environment when handling and processing the samples. These involve decontamination of all work surfaces with the previously mentioned products and sterilization of required material. The worker should wear a clean lab coat and latex gloves that cover all exposed skin that gets in contact with the samples. It is important that surfaces that are in direct contact with the samples are not touched, such as tube taps, hubs of syringes and needle tips.

RESULTS AND DISCUSSION

Harvesting and isolation was achieved for the 10 horses with no restrictions using the previously described technique. The total number of BMMNCs (mean ± SD) and percentage of cell viability observed was 190±54 x 10^6 and 98±1.2%, respectively (table 2). According to Gram staining technique, no evidence of bacterial contamination was observed.

MSCs in horses are commonly retrieved from BM, adipose tissue and umbilical cord blood samples. The sternum is the most commonly used site for aspiration of mesenchymal stem cells derived from equine bone marrow (Taylor and Clegg 2011) and the 5th sternebra has been reported to be most safe of sites, as it has the largest dorsoventral span and is cranial to the apex of the heart (Kasashima et al 2011). Other sites for BM aspiration in the horse include the tuber coxae, tibia, ribs and humerus (Sellon 2006, Taylor and Clegg 2011). However, in the author’s experience, the sternum should be the site of choice to perform BM aspiration in horses because its safer for the practitioner and the patient, and retrieve a larger amount of sample.

Reported complications of sternal BM sampling in horses include cardiac puncture and nonfatal pneumopericardium (Taylor and Clegg 2011). No complications were observed in the 10 horses of this study. That risk can be minimised

<table>
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<th>Horse</th>
<th>1</th>
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<th>7</th>
<th>8</th>
<th>9</th>
<th>10</th>
<th>Mean</th>
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<tbody>
<tr>
<td>Cell concentration (x 10^6)</td>
<td>272</td>
<td>180</td>
<td>147</td>
<td>84</td>
<td>186</td>
<td>179</td>
<td>256</td>
<td>169</td>
<td>213</td>
<td>215</td>
<td>190</td>
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<tr>
<td>Cell viability (%)</td>
<td>96</td>
<td>97</td>
<td>97</td>
<td>99</td>
<td>97</td>
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<td>98</td>
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</table>

Table 2. Concentration per mL and viability of equine mononuclear cells obtained from bone marrow samples of 10 horses. 
Concentración por mL y viabilidad de células mononucleares obtenidas a partir de muestras de médula ósea de 10 caballos.
by carefully placing the needle at right angles (90º) to the surface of the sternebra and closely monitoring and controlling depth (no more than 2cm) of penetration of the needle into the bone during the procedure (Sellon 2006, Kasashima et al 2011). The authors consider that the use of a Komiyashiki needle with the reported length is unlikely to penetrate the thorax. Infection of BM or subcutaneous tissues is a possible concern after BM collection; however, the risk seems to be minimal if strict sterile technique is maintained, and it has never been observed by the authors. Hemorrhage is a potential complication, especially in horses with coagulation abnormalities such as severe thrombocytopenia. Direct pressure at the puncture site is sufficient to control hemorrhage in most horses (Sellon 2006, Taylor and Clegg 2011). If marrow particles are not observed or if the number of mononucleated cells is low, re-aspiration should be considered. Although viscosity of the sample can be suggestive of marrow origin, the
authors do not consider it to be a criterion to evaluate the origin of the sample.

The addition of anticoagulant and DPBS to the syringes used for BM aspiration helps to prevent clot formation. Although the reviewed literature describes the use of sodium heparin as an anticoagulant for the obtention of BM for cell based therapies (Bourzac et al 2010, Taylor and Clegg 2011, Kasashima et al 2011), the authors could not find reported evidence on the advantage or adversity of its use over sodium citrate for horses.

Although collection of 50-60 mL of BM from the sternum of the horse has been reported (Arnhold et al 2007, Fortier and Smith 2008), the sample might contain blood from peripheral circulation since large volumes are unlikely to be obtained solely from one sternebra. Initial results from a MNCs concentration study from the authors suggest the need to obtain at least 15 mL of BM aspirate when the sample is intended to be used as a BMMNCs (Corrêa et al 2012).

The use of Pasteur’s pipette can be replaced by a sterile disposable IV catheter 14G x 2” attached to a 60 mL sterile plastic syringe. These are commonly used at any large animal clinic and should be available to the equine practitioner.

When layering the sample over Ficoll®, care should be taken to do not mix both components, otherwise, low MNCs concentration can be the consequence, as the Ficoll® layer acts as a density filter.

The authors have performed direct cell counting on a Neubauer® chamber and viability through a dye exclusion test. Epifluorescent microscopy in a digital automated counter (Sperm Class Analyzer®) using ethidium bromide/acidine orange staining were also performed and no differences were observed (data not shown). Data regarding MNCs concentration and cell viability obtained with the described method were corroborated with the results of previous investigations (Bourzac et al 2010).

Any cell isolation method can be affected by bacterial or fungal contamination (Taylor and Clegg 2011). It is usually easily identified in cell culture bottles as cells begin to grow slower, the color of the culture medium changes from red to orange/yellow, and fungal hyphae or bacteria are readily identified under phase-contrast microscopy (Taylor and Clegg 2011). During the isolation process, such alterations are not observed because the procedure does not take more than 1 hour. The use of laminar flow hood to reduce risk of contamination is advised, however it does not ensure septic complications will not happen. Besides, in this project Gram staining, an important tool in microbiology (Beveridge 2001, Su RJ and Pei 2011), resulted negative for all samples suggesting the absence of Gram® and Gram- bacterial contamination. Gram staining plays an important role in the rapid presumptive diagnosis, quality evaluation and antimicrobial management of clinical samples in facilities were no cell culture is available (Su R-J and Pei 2011).

The application of complete BM in tendon and ligament injuries has also been reported as a treatment just after collection, with no septic complications (Herthel 2001). Although not mandatory, the suspension of BMMNCs before injection can be done in cell culture medium which contains antibiotics and anti-fungal, as well the horse can receive antibiotics and cells suspended in autologous plasma.

In 8 cases the BMMNCs were used to treat tendon and/or ligament injuries. Horses treated with samples obtained using this technique did not present any kind of clinical complications. This suggests that an aseptic technique can be an effective method to obtain these cells for tendon and ligament injury treatment. No observations were reported on the obtention and isolation of the BMMNCs.

Although BMMNCs hold great promise for future cell-based therapeutic strategies and indeed are currently being used on the treatment of several types of equine athlete’s and other species musculoskeletal injuries in other species, further studies on the availability to apply the previously described technique to treat joint diseases are required.

The results of this study confirmed that aspiration of BM from the sternum and BMMNCs isolation in the horse using the described technique is a cheap and reliable method to obtain MNCs for cell based therapies to treat equine tendon and ligament injuries.

ACKNOWLEDGEMENTS

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REFERENCES


