Isolation and Characterization of Fetal Adnexa-Derived Mesenchymal Stem Cells from Nili-Ravi Buffalo (Bubalus bubalis)

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ABSTRACT
Mesenchymal stem cells (MSCs) are extremely valuable in veterinary and human medicine due to their potential application in regenerative medicine. The purpose of this study was to isolate, differentiate and characterize bovine MSCs (bMSCs) from fetal adnexa including amniotic fluid (AF) and Wharton’s jelly (WJ) of Nili-Ravi buffalo during the second trimester. After slaughtering of the animals, pregnant uteri (n=3) were retrieved and properly disinfected before bMSC isolation. The cells from AF were isolated by centrifugation at 400g for 10 minutes, while from WJ by enzymatic digestion with trypsin-EDTA (0.05%). The isolated cells were studied for their plastic adherence, phenotype identification, metabolic activity and in vitro differentiation ability. The isolated AF and WJ bMSCs were fibroblast-like cells in their phenotype, adhered to plastic, showed similar metabolic potential and population doubling time (PDT), but proliferative activity was initially higher (P<0.05) in WJ-bMSCs. When appropriately induced, both cell types showed mesodermal differentiation into adipogenic and osteogenic lineages which was further affirmed by immunolocalization of fatty acid-binding protein 4 (FABP4) and osteopontin (OST), respectively. However, image analysis revealed that the osteogenic activity of WJ-bMSCs was significantly higher (P<0.05) than that of AF-bMSCs. MSC surface markers (CD73 and CD90) were also positively expressed by both cell types. The study showed that the fetal adnexa of buffalo is a rich source of MSCs for culture and has robust differentiation capabilities.


INTRODUCTION

Over the past two decades, stem cell (SC) research is at the forefront of biological studies as these cells can regenerate, differentiate, and modulate the immune system; hence, they can be used in different therapeutic applications (de Moraes et al., 2017). The most intensively studied SC are mesenchymal stem cells (MSCs) which can be obtained from fetal annexes by non-invasive protocols. Usually, these tissues are disposed of as medical waste otherwise. Unlike adult SCs (ASCs), these cells have better renewability and immunomodulatory properties (Kumar et al., 2015; Campos et al., 2017), which may be attributed to a close association with the fetus or late activation of senescent genes.

Amniotic fluid (AF) and Wharton’s jelly (WJ) are commonly used fetal annexes for the isolation of MSCs. During pregnancy, AF acts as a shielding liquid around the fetus which is composed of secretions and excretions of the fetus and fetal membranes. It also contains factors for fetal growth and development as well as have a good number of SCs (Dev et al., 2012a). De Coppi et al. (2007) reported that AF-derived cells have phenotypical characteristics of pluripotent SC, hence can differentiate into cells derived from different embryonic layers. It is easy to collect AF at the time of parturition as well as during pregnancy through amniocentesis. The WJ is a gelatinous substance which is present in the umbilical cord (UC) where it surrounds the cord blood vessels. The WJ is composed of collagen fibers, proteoglycans and
myofibroblasts stromal cells which can adhere to plastic and differentiate into many types of cells (Carlin et al., 2006).

Buffalo is an important animal of South Asia and Europe as it provides milk, meat and traction power hence called Black Gold (Bilal et al., 2006). Due to the importance of SCs in regenerative medicine and their transdifferentiation ability into many cell lines, they can be used to address many problems of buffalo. However, the importance lies in the fact of identifying cell isolation sites and further their differentiation properties.

There is a dearth of scientific literature on SCs derived from Nili-Ravi buffalo, specifically on bovine MSCs (bMSCs) derived from fetal adnexa; however, some studies reported the achievement of the richness of SCs and WJ with MSCs in humans (Widowati et al., 2019) and bovines (Yadav et al., 2011; Corradetti et al., 2013). This study on fetal adnexa-derived MSCs from Nili-Ravi buffalo aimed to provide a deep insight into their characterization, proliferation and differentiation capabilities.

**MATERIALS AND METHODS**

**Collection and transportation of samples:** For sampling of AF and WJ from Nili-Ravi buffalo, pregnant uteri (n=3) from the 2nd trimester were obtained from the slaughterhouse of Rawalpindi/Islamabad after approval from Institutional Biosafety and Bioethics Committee (IBC,1771/ORIC). The uteri were transported to Cell Culture Laboratory of Cytogenetics, PMAS-Arid Agriculture University, Rawalpindi, Pakistan, where they were disinfected with 70% isopropanol (Sigma, USA) and ethanol (Sigma, USA).

**Isolation, culturing and passaging of cells:** To collect AF-bMSCs, about 50mL of AF was aspirated and centrifuged at 400g for 10 minutes. After discarding the supernatant, pellet was incubated in defined media (DM) containing Dulbecco’s Modified Eagle’s Medium/Ham F-12 (DMEM/F-12), supplemented with 10% fetal bovine serum (FBS, BioWest, France), 1% antibiotics penicillin/streptomycin (100 units/mL and 100 µg/mL, respectively; BioWest, France), 1% amphotericin B (0.25 µg/mL, BioWest, France) at 37°C in the presence of 5% CO₂.

The WJ samples were obtained in Dulbecco’s phosphate-buffered saline (DPBS) without Ca²⁺ and Mg²⁺. WJ was diced into pieces/ explants (~1-2 mm³) and digested with trypsin-EDTA (0.05% and 0.53mM, respectively) for half an hour and subsequently incubated with DM for five to seven days. Following confluence (~80%), bMSCs were trypsinized for passaging towards the next passage up to passage 2 (P2) which was used in subsequent experiments given below. Throughout the experiment, media was changed twice a week to keep the cells healthy.

**Growth kinetics:** Recovered bMSCs from P2 were seeded at a density of 5x10⁴ cells per well in a 48-well cell culture plate (Corning, USA), and growth kinetics were measured on days 3, 6, 9 and 12. After trypsinization, the population doubling time (PDT) was calculated with the following formula (Lu et al., 2018):

\[
PDT = \frac{\log 2 (t - t_o)}{(\log N_t - \log N_o)}
\]

Here PDT: population doubling time, t: harvesting time, t_o: initial time, N_t: harvested cell number, and N_o: seeded cell number.

**Cell metabolic activity:** The extent of metabolism in recovered bMSCs was measured through 3-(4,5-Dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay. For this purpose, bMSCs (5x10⁴ cells per well) were seeded in a 48-well cell culture plate. On every third day till day 12, bMSCs were treated with 150µL working solution (0.25 µg/mL) of MTT per well, incubated for 2.5h and formed formazan was eluted in 100µL of dimethyl sulfoxide (DMSO). Optical density (OD) analysis was performed at 630nm.

**In vitro mesodermal lineage differentiation assays:** Adipogenic- and osteogenic induction of bMSCs at P3 was carried out in a 24-well cell culture plate (Corning, USA) at a seeding density of 2.5x10⁴ cells per well for confluence as described by Rashid et al., (2021).

A. Adipogenic induction: Adipogenesis in bMSCs was induced in a two-step procedure; first induction (48 hours) and second maintenance (seven days). To induce adipogenesis, an induction media containing DMEM/F-12 supplemented with 10% FBS, 1% antibiotics, IBMX (0.1mM; Sigma, USA), rosiglitazone (10µM; BD Chemicals, Germiston, South-Africa), dexamethasone (0.3 mM; Solarbio, Shanghai, China) and 0.142% insulin (5 µg/ml; Nov, Denmark) was used. For maintenance, adipogenic maintenance media containing DMEM/F-12 supplemented with 1% Ex-cyte (Darmstadt, Germany), 0.142% insulin, 1% antibiotics were used. Subsequently, the cells were fixed with 4% buffered formalin (Sigma, USA), followed by oil red-O (ORO) staining at a concentration of 6:4 with distilled water and observed under the microscope. Then ORO elution was performed with absolute isopropanol and optical density (OD) of eluted stain was measured at 510nm.

The surface area occupied by ORO stain within a cell was estimated with the help of ImageJ software (1.50i, National Institutes of Health, USA) by pruning the “Threshold” function at “Auto” settings.

B. Osteogenic induction: For osteogenesis, bMSCs were incubated for 21 days in an osteogenic induction medium containing α-MEM (Gibco, USA) supplemented with 10% FBS, 1% antibiotics, β-glycerol phosphate (1µM; Sigma, USA), 0.01% dexamethasone (1 mM) and ascorbic acid (50 µM; Sigma, USA). Cells were fixed, followed by staining with Alizarin Red-S (ALZ) and observed for mineral deposition at 10x and 40x. For quantification of ALZ, five areas (Fig. 4 I) per well were photomicrographed and analyzed by ImageJ using green channel under the “RGB stack” option at “Threshold” level 75. The measurements of highlighted
Alkaline Phosphate (ALP) activity of osteogenesis: To access the activity of ALP, bMSC were scratched from cell culture plates and analyzed through a commercially available kit (ELITech Group, France) according to the manufacturer instructions. The ALP concentration was measured by utilizing p-nitrophenyl phosphate as a substrate and normalized to the total protein present in the cell lysate.

Immunocytochemistry (ICC): To localize mesenchymal cell markers (CD73; ab137595 and CD90; ab92574), adipogenic specific marker (FABP4; ab85875) and osteogenic specific marker (osteopontin; ab63856) at P3, bMSCs were seeded at a density of 1×10^5 cells per well in a 6-well cell culture plate preloaded with 6mm sterile coverslips. At the culmination of the induction period, the cells were fixed, treated with 0.3% triton (Triton® X-100), and blocked with goat serum (10%). Then the cells were incubated overnight with primary antibodies followed by incubation with secondary antibodies while nuclear identification was done by counterstaining with 2 μg/mL of DAPI (4′,6-diamidino-2-phenylindole). The coverslips were mounted using an anti-bleach mounting medium (Vecta shield, St. Neots, UK) and were visualized under a fluorescence microscope.

Statistical analysis: Statistical analysis was performed using Minitab® 17.1.0 software. The experiments were conducted in duplicate while three replicates of each experiment parameters of growth kinetics and cell metabolic assay were performed and the results were expressed as mean ± SEM. Data was analyzed by one-way ANOVA for multiple mean comparisons using Tukey’s Honestly Significant test (P<0.05).

In vitro mesodermal differentiation: After adipogenic induction for seven days, bMSCs from both the tissues were successfully differentiated into adipocytes, which was evident from rounded cell morphology and accumulation of lipid in the cytoplasm. The presence of intracytoplasmic lipid droplets was confirmed with ORO staining (Fig. 3-I). Control samples cultured in DM/ control media did not show any change in cell morphology, while faint cytoplasmic/ membrane reactivity with ORO was observed in these cells. ORO concentration per cell of bMSCs induced groups was significantly (P<0.05) higher than their respective control groups, with the highest value in WJ-adipo-induced cells (Fig. 3-IIa). Significantly (P<0.05) higher ORO-stained surface area in adipo-induced bMSCs was recorded than their control groups, with the highest value in AF-adipo-induced bMSCs (Fig. 3-IIb). Immunophenotyping expression of FABP4 was expressed successfully among adipo-induced cells of both the tissues (Fig.5e, f). The presence of FABP4 accumulation was evident around unilocular and multilocular lipid droplets in adipose-induced bMSCs of both the tissues.

Under osteogenic induction, bMSCs differentiated into osteogenic lineage cells as they enlarged and modified their morphology contrary to control cells. Many osteogenic foci were observed which were formed by the accumulation of calcium or other minerals [hydroxylapatite mineral (Ca_{10}(PO_{4})_6(OH)$_2$)]. These minerals were seen as an increase in ALP activity of osteogenesis.
Fig. 2: Growth curve (a) and MTT absorbance value (b) of AF-MSCs and WJ-MSCs of Nili-Ravi buffalo (Bubalus bubalis) expressed as mean ± SEM (P<0.05).

Fig. 3 (I-II): Adipogenic differentiation assay of AF and WJ-derived MSCs isolated from Nili-Ravi buffalo (Bubalus bubalis). (I) Intracytoplasmic lipid droplets appear red with ORO staining in induced AF (b) and WJ (d), while control AF (a) and WJ (c) did not show such cytoplasmic changes, only cell membrane lipids show light red color. (II) Lipid concentration per cell (a) and percent lipid surface area per cell (b) of each tissue, where A and B indicate a statistically significant difference (P<0.05) between the control and induced groups.

secreted by the induced cells and appeared red by ALZ staining. Control cells cultured in DM did not show morphological changes and extracellular hydroxyapatite mineral deposits (Fig. 4-IV). ImageJ analysis of bMSCs of osteo-induced and control cells showed that AF and WJ induced cells had a very high extracellular calcium deposition, with WJ-osteod-induced cells had remarkably high calcium deposits (Fig. 4-II) while in control groups, calcium deposition was scarce. ICC revealed the presence of osteopontin among osteo-induced cells of both the tissues (Fig. 5g, h) and was uniformly distributed throughout the cytoplasm that confirms successful differentiation of bMSCs into the osteogenic lineage. Cellular ALP activity is considered an early marker of osteogenesis, which was assessed in the cellular lysate. The converted cells of both origins exhibited an increase of ALP (U/g of protein) compared to control cells. However, WJ-derived bMSCs had significantly (P<0.05) higher ALP activity as compared to AF-derived bMSCs (Fig. 4-III).

Immunophenotyping: An expression of mesenchymal cell surface markers is a characteristic parameter for detecting bMSCs which substantiates the evidence of the mesenchymal nature of cells. The ICC analysis exposed that bMSCs are positive for mesenchymal markers including CD73 and CD90 (Fig. 5a-d). CD73 was localized near the nuclei, while CD90 was evenly distributed throughout the cell membrane. The intense presence of CD73 seems to be present in the Golgi apparatus and further has to be transported on the cell membrane. The results indicated that both sources provide cells that are similar to MSCs.
**Fig. 4** (I-IV): *In vitro* differentiation assay of AF-MSCs and WJ-MSCs isolated from Nili-Ravi buffalo (*Bubalus bubalis*). (I) Schematic diagram of one well of a 24-well plate. The five small shaded squares on a cross represent the microscopic fields used for ALZ digital quantification. (II) ImageJ analysis of ALZ stained area out of the total microscopic field. Osteo-induced cells, specially WJ-MSCs, showed a massive degree of mineralization (mean±SEM) as compared to control, where a, b and c indicate a statistically significant difference (P<0.05) between the control and induced groups. (III) ALP activity of bMSCs (mean±SEM) which was significantly (P<0.05) the highest in induced WJ-bMSCs followed by induced AF-bMSCs. (IV) Positive staining of extracellular calcium/mineral deposition with ALZ which indicates osteogenic lineage transdifferentiation (IV-AF & IV-WJ: b, d). Control cells maintained fibroblast morphology and stained negative with ALZ (IV-AF & IV-WJ: a, c).

**DISCUSSION**

Fetal adnexa is a safe and readily available source of SCs as compared to conventional sources and is recognized as a highly potential source of bMSCs that yields an enormous number of cells in the subculture (Gugjoo et al., 2019). During this trial at P0, AF showed a high number of non-adherent cells which were removed during media change, however, few rounded cells showed plastic adherence which was also reported in Holstein cow (Rossi et al., 2014). Non-adherent cells having flat morphology come from the amniotic membrane, gastrointestinal, urogenital and respiratory epithelium (Dasgupta and Jain, 2016).

Unlike AF, WJ is a newer tissue to get attention as a source of bMSCs in human and veterinary science (Sreekumar et al., 2014), therefore, there is limited data available on growth kinetics and plasticity of WJ-bMSCs. For isolation of bMSCs from WJ two techniques viz, explant attachment method and enzymatic digestion method are used. The former method provides a higher number of cells with higher expression of pluripotency markers while the latter method is easier and yields cells with a high proliferation rate and purity (Widowati et al., 2019). In this study, both methods were combined to obtain the collective effect. Trypsin enzyme hydrolyzed the collagen fibers (Liu et al., 2018) and released bMSCs. WJ-bMSCs were readily attached to plastic which is as per the principles established by the International Society of Cellular Therapy (ISCT) for the identification of MSCs (Dominici et al., 2006). Like other reports (Corradetti et al., 2014), our findings showed more than 90% viable cells; however, Vidane et al., (2014) have reported less than 90% viability in feline. Uranio et al., (2014) reported very low viability (71.40±17.3%) in late passages (beyond P4) compared to early passages (92.83±3.89%) in canines. Therefore, it is recommended to use MSCs at early passages to obtain maximum viable cells.
The trajectory of the growth curve and population doubling time (PDT) are highly variable parameters among MSCs derived from different sources and at different stages of pregnancy. MSCs mostly showed a logarithmic growth pattern which is in line with our findings (Deng et al., 2018). AF and WJ-bMSCs exhibited very high PDT (79.86±6.45 and 72.64±2.02 hours, respectively) which is comparable to that of AF-MSCs of Holstein cows (3.5±1.6 days 84±38.4 hours) (Rossi et al., 2014) and feline (2.8±3.2 days approximately 67.24±68 days) (Iacono et al., 2012). However, it is higher than already recorded in buffalo amniotic membrane (60±5 hours) (Deng et al., 2018) and AF-MSCs of term pregnancy in cattle (2.69±0.32 days approximately 64.56±7.68 hours) (Corradetti et al., 2013). The difference may be due to different types of tissues and stages of pregnancy.

An initial high metabolic activity was recorded in bMSCs that gradually decreased towards day 12. This finding is in agreement with the amniotic SC of rabbits where an initial wave of proliferation was recorded, followed by a drop in proliferation rate, which rebounded at the end of the experiment (Borghesi et al., 2017). These findings are, however, contrary to data obtained from fibroblast cells of cat amniotic membrane (Vidane et al., 2014) who reported continuous growth. This discrepancy may be due to their use of a fewer number of seeded cells per area. There is no data available to compare the results with bovines.

The bMSCs transdifferentiated into mesodermal adipogenic and osteogenic lineages which is evident from ORO and ALZ staining. Previously, fetal adnexa-derived MSCs in bovine (Deng et al., 2018) and ovine (Lu et al., 2018) have been transformed into these lineages. In this study, the cells were converted into adipocytes at the end of the experiment, however, the period of lipid concentration decreased from fourteen days (Jurek et al., 2020) to seven days, which may be considered an achievement in the rapid conversion of bMSCs into functional adipocytes.

The lipid concentration measured in terms of OD value showed the highest concentration in WJ-adipo-induced cells while the image analysis method showed the highest concentration in AF-adipo-induced cells. This difference may be because the elution method measured the total lipid present in the cells, while ImageJ calculated ORO stained surface area. It can be seen that AF-adipo-induced cells were more flattened and occupied a greater surface area than that of WJ-adipo-induced cells which were relatively spherical. This inconsistency can be resolved by using 3D scanning electron microscopic (SEM) images of both cell types.

Osteogenesis was observed in induced AF-bMSCs by ALZ staining which is at par with previously reported findings in bovines (Jurek et al., 2020). Similarly, induced WJ-bMSCs were also confirmed for osteogenic lineage differentiation which has also been reported in adipose-derived bMSCs (Sreekumar et al., 2014). ALZ quantification (Booysen et al., 2018) of stained cells revealed that a very high concentration of extracellular hydroxylapatite mineral was secreted by WJ-bMSCs suggesting that they possess higher osteogenic differentiation potential than AF-bMSCs (Conconi et al., 2011).

The FABP4, which is considered as an intermediate and late maker of adipogenic differentiation assay, was also detected in both bMSCs which is in agreement with previous findings in human AF-MSCs (Conconi et al., 2011) and bovine AF-MSCs (Sandhu et al., 2017; Jurek et al., 2020). Similarly, in this study osteogenic specific gene (osteopontin) was identified in both induced cells, AF- and WJ-bMSCs, which was also reported in cattle-AF (Rossi et al., 2014). As MSCs must express a positive reaction for mesenchymal-specific genes (Dominici et al., 2006), the results of immunofluorescence suggested that AF- and WJ-bMSCs had a positive expression of these mesenchymal markers.

**Conclusions:** It may be inferred that cells derived from AF and WJ of Nili-Ravi buffalo during second-trimester pregnancy are similar to MSCs by their phenotypic characteristics, in vitro differentiation ability, renewability and plastic adherence, however, WJ-bMSCs exhibited better differentiation potential than AF-bMSCs.

**Authors contribution:** This manuscript is from the Ph.D. thesis of AS. ASQ and MAS conceived the idea and designed the study. AS conducted the experiments. All
authors were involved in data interpretation, write-up, and final approval of the manuscript. We, the authors declare no financial/ intellectual conflict of interest with any person/ company or institution.

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