Multilineage Differentiation Potential of Bone and Cartilage Cells Derived from Explant Culture

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Abstract: To date, mesenchymal stem cells (MSCs) from various tissues have been reported, but the yield and differentiation potential of different tissue-derived MSCs is still not clear. This study was undertaken in an attempt to investigate the multilineage stem cell potential of bone and cartilage explant cultures in comparison with bone marrow derived mesenchymal stem cells (BMSCs). The results showed that the surface antigen expression of tissue-derived cells was consistent with that of mesenchymal stem cells, such as lacking the hematopoietic and common leukocyte markers (CD34, CD45) while expressing markers related to adhesion (CD29, CD166) and stem cells (CD90, CD105). The tissue-derived cells were able to differentiate into osteoblast, chondrocyte and adipocyte lineage pathways when stimulated in the appropriate differentiating conditions. However, compared with BMSCs, tissue-derived cells showed less capacity for multilineage differentiation when the level of differentiation was assessed in monolayer culture by analysing the expression of tissue-specific genes by reverse transcription polymerase chain reaction (RT-PCR) and histology. In high density pellet cultures, tissue-derived cells were able to differentiate into chondrocytes, expressing chondrocyte markers such as proteoglycans, type II collagen and aggrecan. Taken together, these results indicate that cells derived from tissue explant cultures reserved certain degree of differentiation properties of MSCs in vitro.

Key Words: Mesenchymal stem cells, cell differentiation, plasticity, multipotent, osteoblast, chondrocyte, adipocyte.

INTRODUCTION

Multipotent mesenchymal stem cells (MSCs) provide a fascinating model of differentiation [1, 2], as well as a promising tool for cell therapy and treatment of a range of degenerative diseases [3, 4]. MSCs are considered to be quiescent, lacking tissue-specific characteristics and maintain this undifferentiated phenotype until they are exposed to appropriate signals [5, 6]. MSCs have a capacity for extensive self renewal and maintain themselves throughout the life of an organism. They are “plastic” and under the influence of specific biological signals differentiate into specialized cells with a phenotype that is completely different from that of the precursor [5, 7]. It is this property that MSCs can be exploited in multiple tissue repair and regeneration through therapeutic cell delivery.

Human bone marrow primarily serves as a reservoir for different populations of stem cells. It contains at least two types of stem cells; hematopoietic stem cells and mesenchymal stem cells (BMSCs) [8-16]. Despite the fact that bone marrow represents the main available source of MSCs, MSCs have been recently identified in numerous other mesenchymal tissues including trabecular bone [17-20], periosteum [21-24], adipose tissue [25-27], synovium [28, 29], skeletal muscle [30], lung [31] and deciduous teeth [32].

These tissue-derived cells have the capacity to differentiate into connective tissue cell lineages, including bone, fat, cartilage and muscle when placed in appropriate environments in vitro and in vivo [11, 33-37]. However, it’s impossible to draw a definite conclusion regarding the differentiation potential of tissue-derived cells due to the different methods used for cell isolation. Even though there is no specific cell surface marker for MSCs, a number of studies selected cell populations from tissue cultures based on the expression of Stro-1, CD105 and CD 106 [38, 39]. Therefore, the current study was undertaken in an attempt to compare the multilineage stem cell potential of tissue explant cultures of bone and cartilage. Bone marrow-derived MSCs (BMSCs) were used as a control and the cells derived from tissue explants of trabecular bone (hOB), cartilage (YCC) and alveolar bone (OBM) tissues were investigated side-by-side as sources of MSCs. Our studies revealed that MSCs isolated from tissue explant cultures had the potential to differentiate into multiple lineages, thereby indicating that these tissues may be alternative sources of mesenchymal stem cells via explant culture.

MATERIALS AND METHODS

Isolation and Culture of Bone Marrow-Derived MSCs

Fresh bone marrow samples (n=7, aged 65-70 years old) were collected by aspiration from femoral canal during knee replacement operation at the Holy Spirit Northside Hospital, Brisbane QLD, Australia, in accordance with the institu-
tional review board guidelines approved by the human research ethics committee of The Queensland University of Technology, The Prince Charles and The Holy Spirit Private Hospitals, Brisbane, QLD, Australia. After draining of synovial fluid, a femoral drill hole was made through the distal femur and an intramedullary rod was passed into the femoral canal. Marrow aspirates were collected by syringe. Mononuclear cells, which include the MSC population, were harvested using LymphoPrep® density gradient centrifugation. Briefly, 2-5 ml of bone marrow was collected in a 50 ml tube containing 5 ml phosphate buffered saline (PBS) supplemented with 200 U/ml Heparin (Invitrogen, Mount Waverley, VIC, Australia). Each sample was filtered through a 100 μm filter and 30 ml PBS was added to the filtrate. The diluted cell suspension was carefully layered below LymphoPrep® and was then centrifuged for 35 minutes at 400 g, at 20 °C without braking. The upper layer was aspirated leaving the mononuclear cell layer undisturbed at the interphase. The MSC-enriched low density fraction was collected carefully and rinsed with 20 ml of control medium containing low glucose (LG; 1g/l glucose) Dulbecco’s-Modified Eagles Media (DMEM; Invitrogen) with 10% fetal bovine serum (FBS; HyClone, Logan, Utah, USA), 10 U/ml penicillin and 10 μg/ml streptomycin (Invitrogen) by centrifugation at 200 g for 10 minutes at 20 °C. The resulting cell pellet was resuspended in DMEM and plated at 8 x 10^5 nucleated cells/T25 cm² tissue culture flasks. MSC cultures were maintained at 37 °C in 5% CO₂ for 6 days before the medium was changed and thereafter the medium was changed every three days. When the cultures were 80-90% confluent, the cells were detached using 0.05% Trypsin-EDTA (Invitrogen) and resuspended at a density of 1.28 x 10^6 cells/T75 flask (termed passage 1) in low glucose DMEM containing 10% FBS and Penicillin/Streptomycin. When the MSC cultures were 80-90% confluent they were replated at a ratio of 1:3 for expansion under the same culture conditions and cryo-preserved in liquid nitrogen for later analysis.

Isolation of Trabecular Bone and Cartilage-Derived Cells

Samples of trabecular bone (hOB) and cartilage (YCC) (n=7, aged 65-70 years old) from the relatively healthy part of tibia of the same patients of marrow collection were obtained. Normal human alveolar bone specimens (OBM) (n=5) were obtained from young, healthy patients aged from 12-16 years old at orthodontic clinic with ethics approval from the University of Queensland. Bone and cartilage pieces were cut into fragments of size 3-8 mm, cleaned of adherent connective tissue, rinsed well and placed directly into six-well culture plates containing high glucose (HG; 4.5g/l glucose) DMEM (Invitrogen) supplemented with 10% FCS (HyClone), 10 U/ml penicillin and 10 μg/ml streptomycin (Invitrogen). Confluent monolayers of fibroblast-like cells were obtained after 10-20 days and were passaged once in 25 cm² flasks until confluent. The cells were then cryopreserved for later analysis. All cultures from individual donors were maintained separately.

FACS Analysis

Analysis of cell surface molecules was performed following the procedure reported by Pittenger [14] and Majumdar [40]. Briefly cells (passage 5) were trypsinized, washed with PBS and in FACS buffer (2% BSA, 0.1% sodium azide in PBS) and aliquots of cells (1 X 10^6) were incubated at 4 °C for 30 minutes with antibodies against CD29, CD34, CD45, CD90, CD105 and CD166 (BD Biosciences, North Ryde, NSW, Australia). Subsequently, secondary antibody IgG1 (fluorochrome-conjugated) was added and incubated at room temperature for 15 minutes. Analysis of the cell surface antigens was performed with a FACS Calibur Cytometer (BD Biosciences, San Jose, CA, USA). Controls for the FACS procedures included conditions where the primary antibody was omitted. In addition, an anti-human IgG1 antibody (BD Biosciences) was used.

Osteogenic, Chondrogenic and Adipogenic Differentiation

Differentiation experiments were performed on cultures recovered from frozen stocks at passages 2-6. The cells were grown to confluence and then subjected to specific differentiating conditions as described below. The differentiation assay was carried out in triplicate in six-well culture plates with all samples of BMSCs (n=7), hOB (n=7), YCC (n=7) and OBM (n=5).

Osteogenic Differentiation

Osteogenic differentiation of subconfluent MSCs from bone marrow and explant tissue-derived monolayers was induced by transfer to high glucose DMEM (HG DMEM) medium containing 10% FBS, 50 μM ascorbic acid 2-phosphate, 10 mM β-glycerol phosphate and 100 nM dexamethasone (Sigma Aldrich, Castle Hill, NSW, Australia) in HG DMEM. After four weeks, calcium phosphate mineral deposition was visualized by von Kossa staining.

Chondrogenic Differentiation

Chondrogenic medium consists of serum-free medium consisting of HG DMEM supplemented with 10 ng/ml TGF–β3 (Chemicon, Boronia, VIC, Australia), 100 nM dexamethasone (Sigma Aldrich), 50 μg/ml ascorbic acid 2-phosphate (Sigma Aldrich), 100 μg/ml sodium pyruvate (Invitrogen), 40 μg/ml proline (Sigma Aldrich) and ITS-plus (BD Biosciences; final concentrations: 6.25 μg/ml insulin, 6.25 μg/ml transferrin, 6.25 μg/ml selenous acid, 5.33 μg/ml linoleic acid and 1.25 mg/ml bovine serum albumin). After four weeks, the differentiated monolayers were stained with Alcian Blue (Sigma Aldrich) for visualization of proteoglycan deposition.

Cells were also cultured as a micromass pellet of cells, created by gentle centrifugation [36]. To initiate the chondrogenic culture, aliquots of 5 x 10^5 cells were suspended in 0.5 ml of chondrogenic media with TGF–β3 (with the rest of the supplements described above) and centrifuged at 600 g for 5 mins in a 15 ml polypropylene conical tube. The cells were left at the bottom of the tube and placed in a 37 °C CO₂ incubator with loosened caps permitting gas exchange. The medium was replaced three times a week. After 3 weeks the cell pellets were washed in PBS, fixed in 4% paraformaldehyde and embedded in paraffin. The pellets were sectioned (5 μm thickness) and analyzed for the expression of proteoglycans using Alcian Blue stain. Immunohistochemical studies were carried out on the pellet sections to detect chondrocyte specific proteins such as aggrecan (AGG) and type II collagen (Col-II).
Adipogenic Differentiation

Cells from passages 2-6 were treated for 3 days with adipogenic induction medium containing 0.5 mM isobutylmethyloxanthine (Sigma Aldrich), 100-200 μM indomethacin (Sigma Aldrich), 1 μM dexamethasone, 10 μg/ml insulin (Invitrogen) in HG DMEM with 10% FBS, and then the medium was changed to adipogenic maintenance medium (HG DMEM with FBS and insulin) for 24 hrs. After the three cycle schedule of 3 days in inducing medium and 1 day in maintenance medium, cells were grown in adipogenic maintenance medium for another 7 days, with the change of medium every three days. Cells were then washed with PBS, fixed with 4% paraformaldehyde and stained with Oil Red O (Sigma Aldrich) to detect lipid.

Histological and Immunocytochemical Analysis

Histological Analysis

The osteogenic-differentiated cells were fixed initially with 4% paraformaldehyde in PBS at pH 7.4 for 15 minutes. The mineralized matrix was evaluated by von Kossa staining. Cartilage matrix deposition by immunocytochemistry was blocked by 10% swine serum (DakoCytomation, Botany, NSW, Australia) in PBS containing 0.1% bovine serum albumin (BSA; Sigma Aldrich) and sodium azide (NaN₃; Sigma Aldrich). After 30 min, sections were incubated with 3% hydrogen peroxide (H₂O₂) for 15 min to eliminate endogenous peroxidase activity. Non-specific protein binding in the sections was blocked by 10% swine serum (DakoCytomation, Botany, NSW, Australia) in PBS containing 0.1% bovine serum albumin (BSA; Sigma Aldrich) and sodium azide (NaN₃; Sigma Aldrich). Sections were then counterstained with Mayer’s hematoxylin (Sigma Aldrich) substrate for 3 minutes. The reaction was stopped by immersion and rinsing the sections in PBS. The sections were then counterstained with Mayer’s hematoxylin (Sigma Aldrich) for 15 seconds and were rinsed in tap water for 5 minutes. Finally, the sections were dehydrated in ascending concentrations of ethanol solutions, cleared in xylene and mounted with coverslips using DePeX mounting medium (Labtek, Brendale, QLD, Australia).

Immunocytochemical Analysis

Chondrogenic differentiation was also evaluated by 3-dimensional pellet culture. The cell pellets were fixed in 4% paraformaldehyde and then embedded in paraffin blocks. Blocks were cut into 5-μm sections. For immunocytochemical analysis, cell sections were incubated with 3% hydrogen peroxide (H₂O₂) for 15 min to eliminate endogenous peroxidase activity. Non-specific protein binding in the sections was blocked by 10% swine serum (DakoCytomation, Botany, NSW, Australia) in PBS containing 0.1% bovine serum albumin (BSA; Sigma Aldrich) and sodium azide (NaN₃; Sigma Aldrich). Sections were incubated with optimal dilution (1: 10) of primary antibodies for collagen type II and aggrecan (Chemicon) overnight at 4 °C. Sections were then incubated with a biotinylated swine–anti-mouse, rabbit, goat secondary antibody (DAKO Multilink; DakoCytomation) for 15 minutes and then incubated with horse radish peroxidase-conjugated avidin-biotin complex (DAKO Multilink; DakoCytomation) for 15 minutes. Antibody complexes were visualized after the addition of a buffered diaminobenzidine (Sigma Aldrich) substrate for 3 minutes. The reaction was stopped by immersion and rinsing the sections in PBS. The sections were then counterstained with Mayer’s hematoxylin (Sigma Aldrich) for 15 seconds and were rinsed in tap water for 5 minutes. Finally, the sections were dehydrated in ascending concentrations of ethanol solutions, cleared in xylene and mounted with coverslips using DePeX mounting medium (Labtek, Brendale, QLD, Australia).

Total RNA Isolation and Reverse Transcriptase-Polymerase Chain Reaction (RT-PCR) Analysis

Total RNA was extracted from 28 day cultures (cell monolayers and 3-D pellets) using Tri Reagent (Sigma Aldrich) as per the manufacturer’s protocol and total RNA concentration was determined spectrophotometrically. The mRNA was reverse transcribed to first strand cDNA from 1 μg RNA for a 20 μl reverse transcription reaction using SuperScript III reverse transcriptase (200 units/μl, Invitrogen) with Oligo d’T as the primer. Expression of the specific mRNAs relative to the osteogenic lineages [alkaline phosphatase (ALP), type I Collagen (Col-I), osteopontin (OPN), osteocalcin (OCN)], chondrogenic lineages [type II collagen (Col-II), and aggrecan (AGG)] and adipogenic lineages [peroxisome proliferator-activated receptor 2 (PPAR2), fatty acid binding protein (AP2)] were examined by RT-PCR. Beta-2 Microglobulin (β2 M) was used as a control for the RT-PCR analysis. PCR was performed using 2 μl cDNA, 1 U of Red Taq DNA polymerase (Sigma Aldrich) and 10 pM of each primer (Table 1) in a 20 μl reaction mixture for 31-35 cycles to make sure

Table 1. Primers for RT-PCR

<table>
<thead>
<tr>
<th>Primers</th>
<th>Forward</th>
<th>Reverse</th>
<th>Product Size</th>
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<td>ALP</td>
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<td>5'TTGTAACGCTTTGAGAGGCC 3'</td>
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<td>Col- I</td>
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<td>5'GTTGGCTTCCTGGTG 3'</td>
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<tr>
<td>OPN</td>
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<td>5'GTTGATGTCCTCTGGTA 3'</td>
<td>347</td>
</tr>
<tr>
<td>OCN</td>
<td>5'ACCCGAGACACCATGAG 3'</td>
<td>5'TGGAGAGGACGAACTG 3'</td>
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<td><strong>Chondrogenic Primers:</strong></td>
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<tr>
<td>Col- II</td>
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<td>5'CAGCACCTGTCTCACC 3'</td>
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<td>AGG</td>
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that amplification was in the linear range for each primer pair. Each cycle consisted of 94 °C of denaturation for 1 minute, 53 °C of annealing for 30 seconds and 72 °C of extension for 30 seconds. Following PCR, the amplified lineage specific transcripts were electrophoresed in 2% agarose gel to enable visualization of the PCR products. The relative density of PCR products was measured using NIH Image V1.62 to analyze one dimensional electrophoretic gel and the expression level of each gene was calculated relative to beta2-microglobulin.

Statistical Analysis

Unless otherwise stated, all results were based on data generated from triplicate studies of each sample of seven BMSCs, hOB, and YCC samples, and five OBM samples. Statistics were analyzed using one-way ANOVA and SNK-q statistical tests from the statistical package SPSS v14 (Chicago, IL). The significant difference was considered at p<0.05.

RESULTS

Growth and Morphology of BMSC and Tissue-Derived Cells

BMSC fractions from density gradient centrifugation failed to spread, migrate or proliferate when cultured in high glucose media (HG DMEM) containing 10% FBS. However, when cultured in low glucose media (LG DMEM), numerous fibroblast-like cells were observed in the culture. Therefore, BMSC fractions from density gradient centrifugation were cultured in low glucose media (LG DMEM) containing 10%FC and numerous fibroblast-like cells were observed after 7 days in the culture. Subsequently, these cells formed colonies which expanded to form MSC-like fibroblastoid cells occupying the entire plastic surface of the culture vessel. Initially, the isolated adherent BMSCs were observed as heterogeneous groups of cells and later became homogeneous as the cells continued to proliferate based on cell morphology. The primary cultures were maintained for 12-16 days before they were detached by trypsinization and subcultured. Cells from the second passage (P2) demonstrated a fibroblast-like, spindle-shaped morphology when viewed under phase contrast microscopy (Fig 1A). All BMSC samples could be expanded over 24 population doublings (8 passages), but in the 8th passage (P8), the spindle or bipolar-shaped BMSCs began to display a broadened, flat morphology. The cells also showed a distinctive swirly pattern when confluent that was consistent with previous findings [41]. About 95-98% of the isolated BMSCs were phenotypically homogeneous when confluent, as determined by their fibroblastic shape. The results indicate that all of these bone marrow-derived cells have morphologic characteristics of MSCs.

Explant cultures from trabecular bone, alveolar bone and cartilage were maintained in HG DMEM supplemented with 10% FCS and penicillin/streptomycin (Invitrogen). After one week of cultivation, hOBs, YCCs and OBMs also yielded numerous fibroblast-like cells (as shown in Fig. 1B, C, D) that subsequently expanded to occupy the whole plastic surface of the culture vessel. Phase contrast microscopy also revealed similar BMSC-like swirly patterns of cell growth among the tissue derived-cells (data not shown). All these tissue-derived cells were able to be explanted over 24 population doubling (8 passages). Furthermore, in contrast to other cell types, OBMs exhibited rapid proliferative potential and became 95% confluent within 3-4 days of passage (1:3 split) when expanded in high glucose media, while other

Fig. (1). Morphology of primary cells cultured in their respective media for 2 weeks. (A) Bone marrow-derived MSCs in low glucose DMEM isolated by density gradient centrifugation; and (B) trabecular bone chip-derived cells (hOBs), (C) cartilage-derived cells (YCCs) and (D) alveolar bone-derived cells (OBMs) in high glucose DMEM (magnification = 200 times).
types of cells required 8-10 days to reach 95% confluence after passage (1:3 split).

**FACS Analysis**

Immunophenotype analysis of BMSC and explant-derived cells was determined using FACS. Flow cytometric analyses demonstrated that BMSCs and explant-derived cells similarly expressed typical MSC specific antigens CD29, CD90, CD105, CD166, and were negative for the early hematopoietic marker and lymphocytic markers CD34, CD45 respectively (Fig. 2).

**In Vitro Differentiation Studies of BMSCs and the Tissue-Derived Cells into Osteoblasts, Chondrocytes and Adipocytes**

To investigate and establish the differentiation potential, the passage 2 and passage 6 of various tissue-derived cells and BMSCs were placed at a density of 2.7 X 10^4 cells/ml in a six-well plate and cultured under appropriate conditions to induce lineage specific differentiation. Bone-derived, cartilage-derived and alveolar bone-derived explant cultures were assayed in parallel. For osteogenic differentiation, cells were cultured in monolayers in the presence of osteogenic supplements and acquired an osteoblastic morphology with the deposition of mineralized extracellular matrix as demonstrated with von Kossa staining (Fig. 3). No mineralized matrix could be detected in the undifferentiated cultures, which were maintained in culture medium alone. Within 2 – 4 weeks BMSCs showed gradual deposition of calcium nodule formation in the cell culture, which covered the entire vessel surface after four weeks. All tissue-derived cells exhibited an osteoblast phenotype with the detectable matrix mineralization. However, the bone nodule size and number in tissue-derived cells were much less than BMSCs in osteogenic culture. The osteoblastic phenotype was further assessed by measuring the expression of alkaline phosphatase (ALP), collagen I (Col I), osteopontin (OPN) and osteocalcin (OCN) using RT-PCR (Fig. 4A) in all samples. All these osteogenic markers were detectable in all four cell types during 2-4 week osteogenic culture. Variation was detected in the mRNA expression level of these markers in each individual sample. For example in the sample presented in Fig. (4A), YCC showed less mRNA expression of Col-I and OPN and BMSCs showed less expression of OCN. However, no statistically significant difference was found in these gene expression amount BMSC, hOB, YCC and OBM in 28 days of osteogenic induction, when all samples (n=7) were summarized and normalized with the house keeping gene (β2M) (p>0.05) (Fig. 4).

![Fig. (2). Analysis of different cell preparations of MSCs, hOBs, OBMs and YCCs for the expression of surface antigens by flow cytometry. Cells were cultured for 5 passages, harvested and labelled with antibodies against human antigens CD166, CD105, CD90, CD45, CD34 and CD29 indicating both positive for CD 166, CD105, CD29, CD90 and negative for CD34 and CD45.](image-url)
Chondrogenic differentiation in monolayers was promoted in early passages of all samples subjected to serum-free chondrogenic medium containing TGF-β3. Under this condition the tissue-derived cells and the BMSCs lost some of their fibroblastic morphology and acquired the polygonal or cuboidal morphology of a typical chondrocyte (Fig. 3). YCC, along with the BMSCs, exhibited a typical chondrocyte phenotype, but hOB and OBM showed less polygonal morphological changes in 4 weeks chondrogenic culture (Fig. 3). After 28 days the chondrocytic phenotype was further confirmed by the expression of collagen type II (Col II) and aggrecan (AGG) using RT-PCR (Fig. 4B) in all differentiated BMSCs and tissue-derived cells, with no expression in the controls (data not shown). A significantly lower expression of Col-II and AGG in OBM was observed in comparison to YCC (p<0.05).

Chondrogenic differentiation was further evaluated in micromass culture for 3-4 weeks in serum-free chondrogenic medium supplemented with TGFβ3. Sections (5 μm) of 3-week old cultures were processed for histological analysis. Within the sections, depositions of sulphated proteoglycans were stained blue by Alcian Blue (Fig. 5A). Immunocytochemical studies also revealed that the major components of the extracellular matrix (ECM) in the chondrogenic micromass cultures were Col II and AGG as shown by the brown color in the sections (Fig. 5A). RT-PCR results also confirmed the expression of Col II and AGG mRNA as observed in immunocytochemical analysis. There was less expression of Col-II and AGG in BMSCs and hOB pellet cultures compared with YCC (Fig. 5B). However, no significant difference in both Col-II and AGG expression amongst BMSC, hOB and YCC in chondrogenic micromass culture, when all samples (n=7) were summarized and normalized to the house keeping gene (β2M) (p>0.05) (Fig. 5).

The adipogenic potential of early passages of BMSCs and explant tissue-derived cells was assayed after inducing adipogenic differentiation. After 4 week of induction, the cell morphology changed and accumulation of lipid rich vacuoles were observed as visualized by oil red O staining (Fig. 3). OBM showed the lowest level of staining for lipid. However, after 4 weeks of induction the adipocyte specific peroxisome proliferators-activated receptor γ2 (PPARγ2) and fatty acid binding protein (AP2) were determined by RT-PCR (Fig. 4C) and found to be expressed in BMSCs and all tissue-derived cells (Fig. 4).

**DISCUSSION**

Despite the fact that bone marrow aspirates have been considered to be the most accessible and enriched source of MSCs [18] autologous bone marrow procurement has limitations. For reasons that are not apparent, there have been variable problems in isolating BMSCs. The number and proliferative capacity of BMSCs depends on donor age, disease
history and the volume of marrow aspirates obtained [19, 32]. MSCs residing in diverse mature host tissues have recently been recognized [28, 29], hence potentially providing alternative sources of MSCs for both basic research and clinical application. In view of this, we compared and characterized tissue-derived (trabecular bone, cartilage and alveolar bone) MSCs using explant culture method with the properties of bone marrow-derived MSCs to evaluate the differential potential of these cells. This study demonstrated that the MSCs harvested from explant culture of committed tissues shared a commonality with bone marrow-derived cells in terms of morphology and immunophenotype. The expression of molecular differentiation markers and their phenotypic potential to differentiate into multiple mesenchymal cell lineages in vitro were demonstrated in tissue-derived explant cultures. Thus, cells from tissue-derived explant cultures possessed certain degree of typical stem cell properties such as being multipotent and able to self renew.

In our experiments the tissue-derived cells were isolated through explant cultures without treating the tissues with any external enzymatic digestion as reported by others [17]. This explant culture method isolated the cells from cartilage and bone depending on the cell potential to migrate out of the bone and cartilage. These cells showed fibroblastic spindle shape similar to BMSCs and expressed typical MSC cell surface markers, negative for CD45 and CD34 and positive for CD29, CD90, CD166, and CD105 (SH2) [42]. Although it is well known that bone marrow-derived MSCs have great proliferation capacity, in the present study the expansion ability of tissue-derived cells showed similarly to that of BMSCs. Tissue-derived and marrow-derived MSCs were both able to be explanted for at least 24 population doublings.
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The highest self-renewal capacity was noted in OBM, which may in part be explained by the fact that these cells were derived from alveolar bone specimens from patients with a younger age. These data, in turn, suggested that the cells that migrated out of the tissue explants had similar characteristics in culture to bone marrow-derived MSCs.

The multilineage differentiation capacity of the tissue-derived cells was assessed by the phenotypical expression of extracellular matrix and differentiation genes relative to mesenchymal cell lineages: osteogenic, chondrogenic and adipogenic. In general tissue-derived cells were able to be differentiated into osteoblasts, chondrocytes and adipocytes because the extracellular matrix and differentiation genes

(8 passages) with similar kinetics of cell growth. The highest self-renewal capacity was noted in OBM, which may in part be explained by the fact that these cells were derived from alveolar bone specimens from patients with a younger age. These data, in turn, suggested that the cells that migrated out of the tissue explants had similar characteristics in culture to bone marrow-derived MSCs.

**Fig. (5).** Chondrogenesis was determined in micromass pellet cultures using bone marrow-derived MSCs, tissue-derived cells (hOBs and YCCs). Cells (500,000) were subjected to low-speed (600g) centrifugation and the cell mass was cultured in defined chondrogenic media with TGF-β3 for 21 days. Chondrogenesis was evaluated by the accumulation of proteoglycans staining with Alcian Blue (5A, magnification = 200 times). Micromass pellet cultures showing morphological changes were subject to immunohistological analysis with an antibody specific for Col-II and AGG (5A, magnification = 200 times). In control sections there was an absence of such phenotypes. All samples were cultured in pellet format for 28 days in the presence of TGF-β3 prior to recovery of total RNA and PCR analysis. 3D-Chondrogenic differentiation resulted in cells that expressed cartilage specific genes (Col-II and AGG) (Fig. 5B). β-2-microglobulin is used as a control for RT-PCR. The graph shows the relative expression level of each gene in relation to the expression of β-2-microglobulin.
typical for differentiated osteogenic, chondrogenic, and adipogenic cells have been detected. However, histologically greater calcium deposition and hence greater osteogenic differentiation potential was observed with the BMSCs compared to the tissue-derived MSCs as detected by von Kossa stain. Moreover, cells derived from cartilage showed a tendency of less mRNA expression of Col-I and OPN compared with cells derived from bone in the osteogenic condition. Interestingly OBM showed lower differentiation capacity into chondrocytes and adipocytes despite OBM was derived from younger aged patients compared with other tissue derived cells, which indicates that age may not be the only determinant of differentiation capacity of tissue-derived cells. We have also demonstrated that tissue-derived cells can be cultured in high density pellet culture in a serum-free chemically defined medium containing TGF β3; a property shared with BMSCs [36, 43, 44]. The mass of the tissue increased over a period of 3 weeks as shown for the BMSCs. The deposition of cartilage matrix such as proteoglycans was detected in the mass culture of tissue-derived cells. Furthermore, RT-PCR analysis of cells in the micromass pellet revealed the expression of the chondrogenic specific marker genes Col II, and AGG, demonstrating the cartilage phenotype of these pellets in the chondrogenic media. These data regarding mesenchymal differentiation lead us to suggest that tissue-derived cells retain some degree of plasticity and their differentiation capacity is more dependent on their microenvironment and less dependent on their primitive status [32, 45].

In summary, our data indicates that MSCs can be sourced not only from bone marrow, but also from other readily available sources like trabecular bone, alveolar bone and cartilage via explant culture. However, mesenchymal stromal cells isolated from other sources are less potent than bone marrow derived MSCs.

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