Bone Marrow-derived Osteoblasts in the Management of Ovariectomy induced Osteoporosis in Rats

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Abstract

Background: The objective of this study is to assess if infusion of osteoblasts can temporarily reverse osteoporosis in rats.

Methods: Osteoporosis was induced in 20 female Sprague-Dawley rats by performing ovariectomy (OVX) that was carried out at 4 weeks of age. At 3 months a biopsy of the iliac crest was made to assess the bone quality and the same site bone marrow was harvested. From the bone marrow aspirate, MSCs were separated. Osteoblasts were then generated and were characterized using Alizarin red staining. Osteoblasts were injected in the tail vein of 10 rats. Two weeks after the injection of osteoblasts, a second biopsy was done. Animals were euthanized after 8 weeks of osteoblasts infusion by overdose of ketamine mixed with xylazine. The whole femurs and lumbar spine were dissected and the specimens were stored in 2% formalin. The specimens were analyzed using HRpQCT (High-resolution peripheral quantitative computerized tomography (μCT 100, SCANCO Medical AG, Brüttisellen, Switzerland).

Results: In all the 10 animals from which bone aspiration was performed, osteoblasts were cultured and transplanted. Analysis showed that there was significant bone formation at bone sites of distal femur and lumbar spine (P<0.001), with increased number of trabeculae and thickness (P<0.001). Further analysis revealed that there was robust bone formation in the animals that had osteoblasts injection.

Conclusions: This preliminary study indicates that osteoblasts infusion can lead to new bone formation in osteoporosis induced by ovariectomy in rats.

Key Words: Osteoporosis; Ovariectomy; Osteoblasts; Mesenchymal Stimulating Cells (MSCs)

Introduction

Osteoporosis is a disease in which the net loss of bone exceeds bone formation and it occurs in women after estrogen loss in postmenopausal age1-4. Postmenopausal osteoporosis (PMO) is a major public health epidemic world over. The importance of PMO is very clear as with expected increased aging of the population the complications such as the hip fractures will treble in 20505-15. Drug therapy for osteoporosis is effective but comes with side effects like any other drug, the effects ranging from simple gastric irritation, myalgias, arthralgias, hypocalcemia, osteonecrosis of jaw to serious infections6-12. At present MSCs are the most widely used stem cells in the research but in vitro expanded MSCs have a short lifetime after in vivo administration. Moreover, the adverse effects of MSCs, especially in the context of tumor modulation and spontaneous malignant transformation makes it difficult to use them routinely13. Inspite of this, many trials are ongoing using MSCs in various conditions.

Previously, we successfully used osteoblasts in the treatment of avascular necrosis of femur in humans14 and in non-union femurs in animals15 and this encouraged us to use osteoblasts in an attempt to treat osteoporosis in ovariectomised (OVX) rats, as they present an ideal preclinical animal model that shows changes due to estrogen deficit, very similar to human skeleton16.

Methods

Osteoporosis was induced in 20 female Sprague-Dawley rats by performing ovariectomy at 4 weeks of age. After obtaining the ethical approval from the Institutional Review Board of Imam AbdulRahman Bin Faisal University, Dammam, Saudi Arabia (Vide number 2015115/2015), 20 Sprague-Dawley female rats were procured and kept for three days before the study was started. All animals were housed and handled in accordance with the guidelines. Animals were kept in large cages with free mobility and fed with standard diet. They were provided with food, water ad libitum and maintained at 25-28 degrees Centigrade. At 3 months, a biopsy of the iliac crest was made to assess the bone quality and from the same site bone marrow was harvested later. From the bone marrow aspirate, MSCs were separated as described by Piao et al. (2005)11.

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The cell suspension was mixed together and centrifuged at 1,000 g for 4 minutes at 37°C. The supernatant and adipose tissue was removed. The cell suspension was transferred to a 15-mL centrifuge tube containing 5 mL of Percoll (1.073 g/mL, Sigma Corp., St. Louis, Missouri, USA). Cells were dispersed by pipetting again and centrifuged at 5,000 g for 30 minutes. The mononuclear cells in the middle layer were obtained, washed three times with phosphate buffered saline (PBS) and then suspended in low-glucose Dulbecco’s modified Eagle’s medium (L-DMEM; Invitrogen, UK) with 20% heat-inactivated fetal bovine serum (FBS; Gibco BRL, Gaithersburg, MD, USA), 100 μM penicillin G, and 100 μg/mL streptomycin. We opted for acceptable methods of marker analysis like Reverse transcriptase PCR (RT-PCR). We analyzed the hMSC cell populations for the expression of CD44, CD90 and CD45 using RT-PCR. Our hMSC populations showed positive amplification for CD44 (+); CD90 (+) and expected negative expression of CD45 (-). These results confirmed that our cell population was MSC. To confirm the osteogenic potential of the MSCs used, BALB/c MSCs (2 x 10⁴ cells/cm²) cells were incubated in CEM until a confluent layer was achieved and then osteogenic medium was added, containing IMDM supplemented with 9 % FBS, 9 % HS, 2 mM L-glutamine, 100 U/mL penicillin, 100 μg/mL streptomycin, 50 ng/mL L-thyroxine (Sigma Aldrich), 20 mM β -glycerol phosphate, (Sigma Aldrich), 100 mM dexamethasone (Sigma Aldrich) and 50 μM ascorbic acid (Sigma Aldrich). Medium was changed every 5th day. The osteogenic differentiation process was followed as per the recommendations of the commercial media manufacturers. The MSC cells were incubated in osteogenic medium for 14 days. Two numbers of 3.5 mm cell culture dishes were also seeded with MSC and were subjected to the same process of osteogenic differentiation parellely. At the end of the differentiation one of the 3.5 mm dish was used for total RNA harvesting (which was later used for RT PCR confirmation of expression of osteopontin marker which confirms the final maturation into osteoblasts. Second dish was used for alizarin red staining which stains into dark orange color when calcium mineralization is seen around the cells (confirms the osteogenic differentiation). The calcium mineralization intensity and distribution under the field of microscope was used roughly to estimate the percentage of osteogenic differentiation which was estimated to be upto 80%. After 15th day, osteogenic differentiated cells from the MSCs were available for transplantation. Once cells were differentiated, and were characterized using Alizarin red staining. Cells after differentiation have been immediately used for the treatment after being suspended in physiological saline and a backup of the batch of the cells had been frozen in -80 degree celsius before being transferred to liquid nitrogen. As many as 25,000 osteoblasts were calculated and suspended in 500 μl of NSS and infused in the tail vein of 10 rats. Animals were euthanized after 8 weeks of osteoblasts infusion by overdose of ketamine mixed with xylazine. The whole femurs and the control group. Table I and Table II gives the volume and density (Conn.D), Trabecular number (DT-Tb.N) mm, Trabecular thickness (DT-Tb.Th) mm and DT-Tb.Sp: Trabecular separation = marrow thickness (DT-Tb.Sp) mm.Quality Control Calibration tests were carried out every day for the two techniques using an external phantom to detect any potential drift of the instrumentation. There was no drift during the period of the protocol. The same observer analyzed all examinations HR-pQCT.

**Statistical Analysis**

For each sample in the reproducibility study, a coefficient of variation (CV) was calculated as the standard deviation of the three repeated measurements divided by the subject mean. Furthermore, the short-term precision errors were then calculated as root-mean-square (RMS) averages of the precision errors for each of the sample. The statistical level of significance was <0.05.

**Results**

There were no complications of infection or deaths in either the group. In all the 10 animals that the bone aspirate was done, osteoblasts were cultured and were transplanted. Figure 1A shows the HRpQCT picture of the distal femur of the study group. The figure shows large number of trabecular pattern quality and quantity as compared to the control group (Figure 1B). Figure 1C shows the magnification of the distal femur in the study group, which highlights the presence of trabecular pattern. In the control group the section shows large empty spaces and reduced number of trabeculae (Figure 1D). Figures 1E study group and sample six of the control Group showing similar changes. Figures 1F is the Sagittal section of the distal femur in the study group showing the condyles filled with new bone with plenty of trabeculae while Figure 1G is the study group’s sagittal section of the distal femur showing the condyles filled with new bone with plenty of trabeculae compared to control group (Figure 1H) showing wide spaces of no bone to minimal bone in the areas examined when compared to the study group. Figures 9 and 10 are that of HRpQCT scans of the lumbar spine in the study and the control groups show decisively more bone in the study group as compared to the control group. Figures 2 A-D shows the HRpQCT reconstruction of Trabecular Pattern of Vertebral Body in study group. Figures 3A and B shows the histological sections in the study and the control group. Table I and Table II gives the various indices of the Femur and the spine.

**Discussion**

Our study shows that the culture-expanded osteoblasts were effective in increasing bone formation in O VX rats. Both (HRpQCT and doubled labeled tetracycline biopsy) objective assessment of osteoporosis in the two groups gave a promising picture. Secondly our result also shows and we speculate that the effect observed was due to homing of the infused osteoblasts on to the bone surface rather in the other tissues of the body to produce the desired results. Wang et al. [17] used MSCs and reported that these cells increased the strength in the osteoporotic bone, while Ocarino et al. (2010)[18] injected BM MSCS in the bone marrow of osteoporotic rats and concluded that osteoporosis can be treated by injection of BM MSCS. Our strategy was to study the effect of osteoblasts rather MSCs because in an earlier study we found that direct osteoblast infusion at the fracture site gave robust fracture healing[19]. It is accepted now MSCs act via multifaceted pathways, which is not completely understood to increase bone regeneration and homing of the infused cells at the site of intended location[19]. Moreover, difficulty of homing of the MSCs onto the bone surface thereby, low bone forming effect[20]. Recently, Kiernen et al.[21] used minimally expanded exogenous MSCs in a mouse model showed enhanced bone formation. With use of osteoblasts although we observed enhanced new bone formation in the study group of animals, we believe we need to replicate the same in a larger animal to further confirm our findings.
Figure 1: A - Section of Distal Femur of the Study Group showing dense trabeculae; B - Section of the Distal Femur of the Control Group sparse trabecular pattern; C - HRPQCT reconstruction of Trabecular Pattern of Distal Femur in Study Group indicating dense bony structure; D - HRPQCT reconstruction of Trabecular Pattern of Distal Femur in Control Group reveals wide trabecular spaces and empty areas; E - HRPQCT reconstruction of Trabecular Pattern of Distal Femur in Study Group of another animal showing high quantity and quality of the areas; F - HRPQCT reconstruction of Trabecular Pattern of Distal Femur in Control Group of another animal showing sparse bone; G - Sagittal section of the distal femur in the study group showing the condyles filled with new bone with plenty of trabeculae; H - Sagittal section of the distal femur in the control group showing wide spaces of no bone to minimal bone in the areas examined when compared to the study group.

Figure 2: A - Section of Vertebral body of the Study Group showing the area examined with good quantity of the trabecular pattern; B - Section of Vertebral body of the Control Group with empty areas and thinned out trabeculae; C - HRPQCT reconstruction of Trabecular Pattern of Vertebral Body in Study Group depicting the dense trabeculae; D - HRPQCT reconstruction of Trabecular Pattern of Vertebral Body in Control Group with empty spaces when compared to the study group.

Figure 3: A - H & E X400 Section of Distal Femur in Pre injection of Osteoblasts, showing wide area of immature bone and cartilage; B - H & E X400 Section of Distal Femur in Post injection of Osteoblasts after 8 weeks showing formation of new bone in the entire area of the field.
### Table 1: Structural Indices of Distal Femurs

| Parameter                  | Study Group       | Control Group     | P Value |
|----------------------------|-------------------|-------------------|---------|
| VOX-TV                     | 45.04±4.54        | 38.61±1.02        | <0.001  |
| VOX-BV                     | 2.16±0.46         | 1.098±0.22        | <0.001  |
| Connectivity Density       | 16.57±4.5         | 4.92±0.31         | <0.001  |
| Trabecular Number          | 0.7±0.1           | 0.54±0.11         | <0.001  |
| Trabecular Thickness       | 0.07±0.01         | 0.06±0.01         | <0.01   |
| Trabecular Spacing         | 1.40±1.4          | 1.59±0.17         | <0.006  |
| Mean 1 mg/HA/ccm           | 60.69±35.55       | 45.77±20.15       | <0.05   |
| Mean 2 mg/HA/ccm           | 1088.17±8.99      | 1048.10±4.8       | <0.001  |
| TRI – Total Volume         | 42.86±2.70        | 38.21±1.02        | <0.001  |
| TRI – Bone Volume          | 1.74±0.80         | 1.048±0.22        | <0.001  |
| TRI Bone Surface           | 62.89±5.77        | 36.58±5.71        | <0.001  |
| TRI- Trabecular Number     | 0.75±0.33         | 0.48±0.08         | <0.001  |
| TRI- Trabecular Thickness  | 0.05±0.01         | 0.043±0.01        | <0.007  |
| TRI- Trabecular Spacing    | 1.65±0.93         | 2.08±0.39         | <0.006  |

VOX= Based on counting voxels, TRI= based on triangularization of surface.  Mean 1= mean voxel values of everything within volume of interest (Bone and Background)  Mean 2= of segmented region only what was considered bone.

### Table 2: Structural Indices of Vertebral Bodies

| Parameter                  | Study Group       | Control Group     | P Value |
|----------------------------|-------------------|-------------------|---------|
| VOX-Tot Volume             | 38.79±8.03        | 27.9±1.03         | <0.001  |
| VOX-Bone Volume            | 5.33±1.82         | 2.32±0.14         | <0.001  |
| Connectivity Density       | 36.92±4.86        | 15.84±0.72        | <0.001  |
| Trabecular Number          | 1.69±1.31         | 1.14±0.04         | <0.001  |
| Trabecular Thickness       | 0.073±0.01        | 0.05±0.02         | <0.005  |
| Trabecular Spacing         | 0.65±0.05         | 0.82±0.04         | <0.001  |
| Mean 1 mg/HA/ccm           | 196.81±4.65       | 145.68±2.07       | <0.001  |
| Mean 2 mg/HA/ccm           | 1070.32±7.89      | 1032.53±8.99      | <0.001  |
| TRI – Total Volume         | 38.18±7.94        | 26.46±2.94        | <0.001  |
| TRI – Bone Volume          | 5.30±1.84         | 2.22±0.16         | <0.001  |
| TRI-BS                     | 171.14±3.18       | 84.42±0.97        | <0.001  |
| TRI- Trabecular Number     | 2.19±0.23         | 1.49±0.04         | <0.001  |
| TRI- Trabecular Thickness  | 0.06±0.01         | 0.051±0.01        | <0.01   |
| TRI- Trabecular Spacing    | 0.39±0.05         | 0.62±0.02         | <0.001  |

VOX= Based on counting voxels, TRI= based on triangularization of surface.  Mean 1= mean voxel values of everything within volume of interest (Bone and Background)  Mean 2= of segmented region only what was considered bone.
There are two issues, which need to be addressed. How long can we use bisphosphonates for treatment of osteoporosis is still unknown? The long-term effects of drug therapy of osteoporosis are serious and detrimental to the bone itself. Atypical femoral fractures, osteonecrosis of the jaw and safety concerns in patients with renal and gastro-intestinal diseases are well known. The newer drugs like denosumab might ameliorate some of the complications but cannot eradicate them completely. Hence, a search of alternate treatment modality to treat patients with osteoporosis for years with no complications to prevent fractures remains essential. Secondly life expectancy is on the rise and passed over 100 years in some countries and is expected to continue to rise even in the developing countries, making morbidity to rise many fold in patients suffering with osteoporosis.

Our study has some limitations and one such is that we did not perform the homing of the osteoblasts on to the bone surface which could have give strength to the study and differentiate from the MSCs which are suppose to be trapped in the pulmonary capillaries.

This study opens wide array of questions and possibilities. Even though our study shows that in experimental animals osteoporosis can be treated by osteoblasts infusion and it will require more studies to confirm this for a routine use. If the studies do confirm the ability to treat osteoporosis by osteoblasts infusion then how it could be made available for general use. It makes more questions than answers to the issue of alternate treatment of osteoporosis, but stem cell therapy of osteoporosis is a potential treatment modality, which should not be ignored.

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Abbreviations

OVX  - Ovariectomy
MSCs - Mesenchymal stimulating cells
HRpQCT - High resolution peripheral quantitative computerized tomography
PMO - Postmenopausal osteoporosis
TV  - Total volume
BV  - Bone volume
RBV - Relative bone volume
Conn. D - Connectivity density
DT-Tb.N - Trabecular number
DT-Tb.Th - Trabecular thickness
DT-Tb.Sp - Trabecular separation = marrow thickness

Potential Conflicts of Interests

None

Prior Presentations

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