Co-implantation of bone marrow mesenchymal stem cells and chondrocytes increase the viability of chondrocytes in rat osteo-chondral defects

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Abstract. Replacement of chondrocytes by adult stem cells was believed to improve the performance of autologous chondrocytes transplantation, since less chondrocytes were needed. Previous studies have demonstrated that the increased cartilage production in pellet co-cultures of chondrocytes and bone marrow mesenchymal stem cells (BMSCs) is due to the trophic effects of the MSC by stimulating chondrocyte proliferation and matrix production. However, the destination of MSCs or chondrocytes after implanted in osteo-chondral defects is not clear. The aim of the present study is to investigate the viability of MSCs and chondrocytes after co-implantation into a rat osteo-chondral defect model. MSCs were isolated from bone marrow and chondrocytes were extracted from knee joints of neonatal rats. Results of sulfated glycosaminoglycans (GAG) and collagen quantification demonstrated that co-culture pellets of BMSCs and chondrocytes have more GAG deposition than that of BMSCs or chondrocytes alone. Tracking cells with fluorescence protein demonstrated that MSCs disappeared following co-culture. In a rat knee injury model, co-implantation of BMSCs and chondrocytes contained more viable chondrocytes than chondrocytes implanted alone. To conclude, BMSCs were replaced by chondrocytes in pellet co-culture and BMSCs increased the viability of chondrocytes following co-implantation in a osteo-chondral defects model. Co-implantation of BMSCs and chondrocytes may be a promising approach to repairing osteo-chondral defects in the clinical setting.

Introduction

Cartilage is an important connective tissue that exists in the muscle-skeleton system, particularly in joints, rib, ear, nose, bronchial tubes and intervertebral discs (1). It’s generally believed that cartilage contains only one cell type called chondrocyte that produces all cartilage extracellular matrix consisting of Type II collagen in articular cartilage or mixture of type I and type II collagen in fibrocartilage (2). Since cartilage is avascularized, the metabolic activity of chondrocytes is low, compared with other connective tissues (3). Nutrition of chondrocytes is supplied by diffusion, by the pumping action generated by compression of the articular cartilage during extension or flexion of joints (3). Furthermore, chondrocytes are embedded in spaces called lacunae, which keeps them from migrating to damaged areas. Therefore, the self-repair ability of damaged cartilage is limited (4). Over the last decades, stem cell based technology has been proposed for cartilage repair joint injury (reviewed in 5). Especially when bone marrow mesenchymal stem cells (BMSCs) were demonstrated to be able to differentiate into chondrocytes, these were immediately considered as the ideal source for engineering cartilage tissue (5).

However, novel findings uncovered other properties of BMSCs rather than differentiation into specific cell types. One of the most important is the trophic role of BMSCs in tissue repair (4,5). As first introduced, the terminology ‘trophic’ initially referred to non-neurotransmitter bioactive molecules produced by nerve terminals (6). Specifically, ‘trophic’ was first used to describe the process in which BMSCs secrete factors stimulating neighboring cells to release functionally bioactive molecules (7). This term also relates to the effect of the factors produced by BMSC on viability, proliferation, and matrix production of the neighboring cells. This supportive effect of BMSCs on other cells types significantly broadened the application of BMSCs in regenerative medicine. While traditionally it was believed that BMSCs mainly repair damaged tissue by differentiating into specific cell types and replacing lost cells (8), nowadays the trophic role of the BMSCs in tissue repair is considered more important than before (9).

The use of BMSCs to partially replace chondrocytes may reduce the number of chondrocytes necessary for autologous
chondrocytes transplantation (ACT). In previously published papers, pellet co-culture models of chondrocytes and bone marrow derived BMSCs were employed to study the beneficial effects of co-culture on cartilage matrix formation (10-12). In these pellet co-cultures, it had been demonstrated that cartilage matrix was mainly produced by chondrocytes but not BMSCs. These studies revealed a new mechanism of cross-talk between cells in a co-culture model of BMSCs and chondrocytes. Studies indicated that the beneficial effects on cartilage matrix formation in co-culture pellets were due to trophic effects of BMSCs which stimulated chondrocyte proliferation and cartilage matrix deposition. Studies also demonstrated that these trophic effects are independent of culture conditions and BMSCs sources (11). Notably, it’s been identified in all these studies that the ratio of BMSCs decreased dramatically due to BMSCs death and proliferation of chondrocytes in co-culture. However, none of the published studies actually tracked the fate of BMSCs or chondrocytes following co-implantation in an osteo-chondral defect models (6,7,9,10).

In the present study, the destiny of BMSCs and chondrocytes in co-culture pellets and in co-implantation of an osteo-chondral defect model was investigated. Data revealed that BMSCs increased the viability of chondrocytes in co-implantation in an osteo-chondral defect model.

Materials and methods

Cell culture. The use of experimental animals in the present study was approved by the Medical Ethical Committee of the First Affiliated Hospital of Bengbu Medical College (Bengbu, China). Primary chondrocytes were obtained from knee joints of 12 neonatal Sprague Dawley rats, all male, aged 8 weeks. The rats were kept in 12-h light/dark cycle at 37°C and a humidity of 40% and had free access to water and food. The rats were anaesthetized with isoflurane at a dosage of 1-3%, as previously described (13). Cartilage biopsies were digested for 20-22 h in collagenase type II (0.15% Worthington, NJ, US) dissolved in medium containing DMEM supplemented with 10% FBS (Gibco; Thermo Fisher Scientific, Inc., Waltham, MA, USA) and antibiotics (100 U/ml penicillin and 100 µg/ml streptomycin; basic medium) as previously described (14). BMSCs were isolated from the bone marrow of neonatal rats previously reported (13,14). BMSCs were seeded in culture flasks with basic medium. Media were replaced every 2 days to remove floating cells. These experiments were performed in accordance with the recommendations and guidelines for the care and use of laboratory animals established by the Chinese National Committee for the Use of Laboratory Animals. The rats were kept in 12-h light/dark cycle at 37°C and humidity of 40% and had free access to water and food. The rats were anaesthetized with isoflurane with dosage of 1-3% and a 12-h light/dark cycle at 37°C. All reagents used were purchased from Sigma-Aldrich; Merck KGaA, Darmstadt, Germany.

Pellet culture and chondrogenic culture. For mono-cultures, 200,000 primary chondrocytes or BMSCs were seeded in one well of a round bottom 96 wells plate (non-tissue culture treated). For co-cultures, 200,000 chondrocytes or BMSCs were seeded at 1:1 ratio. Cells were initially seeded in basic medium and centrifuged at 37°C for 5 min at 500 x g. Medium was changed to chondrogenic differentiation medium (DMEM supplemented with 40 µg/ml of proline, 50 ug/ml ITS-premix, 50 ug/ml of AsAP, 100 ug/ml of Sodium Pyruvate, 10 ng/ml of TGFβ3, 10⁻² M of dexamethasone, 500 ng/ml of BMP6, 100 U penicillin/ml and 100 µg/ml streptomycin) one day following seeding and stable pellets were formed. Cell pellets were cultured for 4 weeks prior to analysis.

GAG staining. Cell pellets for co-cultures were fixed with 10% formalin for 24 h, decalcified, dehydrated and embedded in paraffin using routine procedures (14). A microtome (Thermo Fischer Scientific, Inc.) was used to cut 5 µm thick sections. Slides were then deparaffinized and stained for sulfated glycosaminoglycans (GAG) with Toluidine blue for 2 h at 37°C.

Quantitative GAG and DNA content assays. Cell pellets were washed with PBS and stored at -80°C for 16-20 h. Subsequently, they were digested with proteinase K solution [1 mg/ml proteinase K in Tris/EDTA buffer (pH 7.6)] for >16 h at 56°C. GAG content was spectrophotometrically determined with 1,9-dimethylmethylen blue chloride (DMMB) staining for 2 h at 37°C in PBE buffer (14.2 g/l Na₄P₂O₇ and 3.72 g/l Na₂EDTA, pH 6.5) using a microplate reader (TECAN group, Ltd., Mannedorf, Switzerland) at an absorbance of 520 nm using standard curves generated with chondroitin sulfate. Total DNA content was determined using a CyQuant DNA Kit (Molecular Probes; Thermo Fisher Scientific, Inc.).

DNA isolation and reverse transcription-quantitative polymerase chain reaction (RT-qPCR). DNA samples from cell pellets were isolated with DNeasy Mini Kit (Qiagen GmbH, Hilden, Germany). Total genomic RNA was used for RT-qPCR using the iQ SYBR Green Supermix (Bio-Rad Laboratories, Inc., Hercules, CA, USA). PCR Reactions were carried out on MyiQ2 Two-Color Real-Time PCR Detection System (Bio-Rad, Laboratories, Inc.) under the following conditions: cDNA was denatured for 5 min at 95°C, followed by 45 cycles, consisting of 15 sec at 95°C, 15 sec 60°C and 30 sec at 72°C. For each reaction, a melting curve was generated to test primer dimer formation and non-specific amplification. Primer sequences were as follows: Green fluorescence protein (GFP) Forward, 5'-ACGACGGCAACTACAAGA CC-3' and Reverse, 5'-TTGTACTCCAGCTTTGTCGCC-3'; red fluorescence protein (RFP) Forward, 5'-AACCTGCAAGTGTACCAAGG-3' and Reverse, 5'-CAATAGTCGGG GATGTCGG-3'; GAPDH Forward, 5'-GATTGTGAGAGG CCGTGTGA-3' and Reverse, 5'-TTCTCAGCTTGGACTGTG CC-3'. Relative gene copies were calculated using the 2⁻ΔΔCq method (15). GAPDH was used for normalization.

Rat osteochondral defect model. Twelve athymic nude rats (all male) of 8-weeks old, weighing 25-30 g, were kept in 12-h light/dark cycle at a temperature of 37°C and a humidity of 40% and had free access to water and food. They were anaesthetized with isoflurane with dosage of 1-3% and a 12-h light/dark cycle at 37°C. All reagents used were purchased from Sigma-Aldrich; Merck KGaA, Darmstadt, Germany.
move freely following surgery. Each rat carried one pellet. Pellets made of BMSCs or chondrocytes, or co-cultures were implanted into the three experimental groups, containing 4 rats per group. All samples were used for histological staining.

**Cell tracking with green and red fluorescence proteins.** To track cells in co-cultures and co-implantation, BMSCs were labeled with red fluorescence protein by lenti-viral transduction, while chondrocytes were labeled with green fluorescence protein. Lenti-GFP and Lenti-RFP virus were purchased from lenti-virus were carried out by Hanbio biotechnology, Co, Ltd. (Shanghai, China). Both lenti-viral vectors contained resistant genes against puromycin. Infection of cells with Lenti-GFP or Lenti-RFP (4%) was performed in 1.0 ml of serum-free basic medium for 4 h at 37°C. Following infection, the remaining supernatant was removed and replaced with basic medium supplemented with 10% fetal bovine serum. Stably transduced cells were selected by adding puromycin (1 µg/ml) in culture medium on day 2 following infection. Antibiotic selection lasted for 1 week. Selection efficiency was verified by visual examination under a fluorescence microscope. To examine GFP and RFP signals in co-culture pellets, cryosections were made with a cryotome (Leica CM1520, Leica Microsystems GmbH, Wetzlar, Germany).

**Immunofluorescence staining.** For immunocytochemistry, sections of co-culture pellets of BMSCs and chondrocytes were deparaffinized, incubated with 3% hydrogen peroxide and blocked in 1% bovine serum albumin and 1.5% normal goat serum at 37°C. Slides were subsequently incubated overnight at 4°C with mouse monoclonal antibodies against GFP (GF28R; Novagen; Merck KGaA) or RFP (cat no. 69831-3; Novagen; Merck KGaA). Sequentially, primary antibodies were visualized by incubating with fluorochrome-labeled secondary antibodies at 37°C (L21998; Invitrogen; Thermo Fisher Scientific, Inc). Counterstaining was performed with DAPI under a light microscope (Olympus IX51; Olympus Corporation, Tokyo, Japan).

**Statistical analysis.** Statistical significance between different groups was examined with one-way analysis of variance followed by Tukey Honestly Significant Difference Test. All data were presented as the mean ± standard deviation. P<0.05 was considered to indicate a statistically significant difference.

**Results**

**Isolation and labeling of rat BMSCs and chondrocytes.** Articular chondrocytes were isolated from knee joints of neonatal rats. Following in vitro expansion for two passages, cells displayed a typical morphology of chondrocytes (Fig. 1). Chondrocytes first spread in a bipolar fashion for a few days, then most cells displayed polygonal morphology with few filopodia. BMSCs presented a spindle-like shape. With time in culture, cells gradually adopted a more fibroblastic morphology (Fig. 1). With lenti-virus infection, chondrocytes were labeled with GFP while BMSCs were labeled with RFP. Both cells were selected by puromycin for 1 week prior to reaching a labeling efficiency closing to 100%, as examined by fluorescent microscope (Fig. 1).

**Co-culture pellets deposited more GAGs and collagen than mono-cultures pellets.** Labeled chondrocytes and BMSCs were used to make cartilage tissue in pellet cultures. Four weeks following seeding, cell pellets of both mono- and co-culture were harvested for histology, GAG assay. As demonstrated in Fig. 2A, BMSCs pellets were able to deposit some GAG into extracellular matrix. Chondrocytes pellets, however, deposited much more GAG than BMSCs. When co-cultured, chondrocytes and BMSCs together produced extracellular matrix containing abundant GAG in the pellets which was significantly more than that in mono-culture pellets, as measured by toluidine blue staining (Fig. 2B). The same was observed in the collagen assay by quantifying hydroxyproline (Fig. 2C).

**BMSCs and chondrocytes in co-cultures.** To track BMSCs and chondrocytes following co-culture, cryosections were made to examine GFP and RFP signals in the pellets, 1 week after the pellets were made. As depicted in Fig. 3A, both RFP and GFP positive cells were present in co-culture pellets. RFP labeled BMSCs tended to present in the center of the pellets, while GFP labeled chondrocytes were distributed more on the periphery of the pellet. Subsequently, RT-qPCR was performed to track RFP and GFP DNA contents in them. Compared with the cells initially seeded (week 0), the GFP positive cells dropped following 1 week of co-culture to roughly 80%, continued to decrease until week 3 and kept stable at week 4 at about 15% (Fig. 3B). On the other hand, GFP positive cells increased at week 1 to about 120% of week 0, and remained stable thereafter (Fig. 3C).

**Effect of BMSCs on the survival rate of chondrocytes following co-implantation in rat osteo-chondral defect model.** To test the viability of cells following implantation into osteo-chondral defects, pellets made of BMSCs, chondrocytes or co-cultures...
were implanted into a nude rat knee injury model. Cell tracking by DNA content using GFP and RFP indicated that a few BMSCs survived in defects when implanted alone (Fig. 4A). In one section, less than 40 cells were found to be red in the defect area (Fig. 4B). Chondrocytes however, survived compared with BMSCs when implanted alone into the defects, with about 80 cells labeled with green in one section. Notably, the viability of BMSCs in co-culture pellets was extremely low, with roughly 4 cells found in defect area (Fig. 4C).

**Discussion**

In recent years, trophic effects of BMSCs have attracted much attention in cartilage engineering (reviewed in 16).
However, very few studies successfully revealed the viability of BMSCs or chondrocyte after co-implantation into osteo-chondral defects (13,15). In the present study, BMSCs and chondrocytes were labeled with fluorescence proteins and were tracked in pellet co-cultures and in a osteo-chondral defect model.

Multiple studies have investigated the effects of BMSCs on chondrocytes (13-15), but few have provided data to indicate the viability of BMSCs and chondrocytes in the content of direct cell-cell contact neither in vitro nor in vivo (13-16). The present study demonstrated that co-culture of BMSCs and chondrocytes led to a decrease in BMSCs cell numbers which can be explained by reduced cell proliferation or cell death, but this needs to be confirmed. Since previous study demonstrated massive cell death of BMSCs by apoptosis (10). The decrease of BMSCs numbers following 1 week of co-culture may suggest that chondrocytes were able to change their behavior. Similar results have been demonstrated in studies using co-culture of BMSCs with chondrocyte pellets from different sources, during 3 to 4 weeks of culture, in which BMSCs numbers decreased progressively (17,18). Besides the dramatic decrease of BMSCs following co-culture, the proliferation of chondrocytes demonstrated a trend of slowdown of the increasing rate following one week of co-culture. In fact, in a previous study employing a pellet co-culture system of BMSCs and chondrocytes, an increase in chondrogenic markers was observed at later stage of culture, following day 7 (19). This is in line with the present finding that proliferation of chondrocytes occurred in the first week of co-culture (20).

A previous report demonstrated that significant numbers of TUNEL positive BMSCs were detected in co-culture pellets (21). This result suggests that the BMSCs may have died by apoptosis upon contacting with chondrocytes. This may have occurred due to cell compaction, and nutrition or space limitation in pellets. The cell labeling experiments demonstrated that the majority of BMSCs reside in the center while most chondrocytes presented on the edge of the pellets. BMSCs may undergo apoptosis simply due to lack of nutrients or oxygen, which was more likely to happen in the center of the pellet (22). However, this is not sufficient to explain why the number of BMSCs deceased so much since some of BMSCs
did survive in the middle of a pellet made by BMSCs alone. This suggested that besides to location of BMSCs in pellets, the presence of chondrocytes may have also contributed to the apoptosis of BMSCs. It’s known that chondrocytes may secrete some apoptotic cytokines (23), which may have induced the death of BMSCs, but his needs to be further confirmed. Furthermore, it is likely that direct cell-cell contact between chondrocytes and BMSCs may have contributed to increased cell death (24).

Viability of cells in tissue engineering is an important issue for regenerative medicine (25). The present data indicated that viability of chondrocytes increased a lot with co-implantation compared with chondrocyte implanted alone, despite the death of BMSCs. A lot of factors are involved in the death of chondrocytes during autologous chondrocyte transplantation into joint environment (26). Mechanical stress, low oxygen and lack of nutrients may cause apoptosis or necrosis of chondrocytes. Small chemicals and bio-compatible scaffolds are designed to increase the viability of chondrocytes (27,28). Findings of the present study may provide an alternative solution to reduce the cell death of chondrocytes after implantation, which is mixing chondrocytes with BMSCs. This may benefit the matrix deposition, but also increase chondrocytes viability, but this needs to be studied further.

To conclude, the present data indicated that BMSCs were overtaken by chondrocytes in the pellet co-culture. Results from the in vivo study demonstrated that BMSCs increased the viability of chondrocytes following implantation in osteo-chondral defects. Co-implantation of BMSCs and chondrocytes may be a promising procedure in repairing osteo-chondral defects in clinical settings.

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Availability of data and materials

The datasets generated and analyzed during the current study are not publicly available due to statutory provisions regarding data and privacy protection, the dataset supporting the conclusions of this article is available upon reasonable request directed to the corresponding author.

Authors' contributions

ZZ, XZ and JG were involved in the conception and design of the study, in the collection, assembly, analysis and interpretation of the data, and in drafting of the article; they also provided statistical expertise. JZ, MW and ZZ contributed to final approval of the article, provision of study materials, and administrative, technical and logistical support, as well as critical revision of the article for important intellectual content.

Ethics approval and consent to participate

The use of experimental animals in the present study was approved by the Medical Ethical Committee of the First Affiliated Hospital of Bengbu Medical College (Bengbu, China).

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

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