Basics of isolation and cultivation of chondrocytes according to good laboratory practice
Principios en el aislamiento y cultivo de condrocitos de acuerdo a buenas prácticas de laboratorio

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Resumen
Objetivos: El objetivo del presente estudio era determinar si los condrocitos aislados de cinco pacientes ancianos (edad media 63 años) con artrosis (grado 3) mantienen su proliferación y potencial condrogénico. El aislamiento y cultivo de condrocitos fueron llevados a cabo de acuerdo a los estándares de buenas prácticas de laboratorio.

Métodos: Los condrocitos fueron aislados de una biopsia de cartílago mediante digestión enzimática. El cultivo fue llevado a cabo en un ambiente controlado (sala blanca). La caracterización del fenotipo de los condrocitos se logró mediante análisis de citometría de flujo.

Resultados: Tras tres semanas de cultivo se podían observar estructuras poligonales propias de los condrocitos, pero también se observaba morfología de tipo fibroblasto en el cultivo. El análisis de la citometría de flujo reveló que el fenotipo de los condrocitos cultivados tras el primer pasaje era positivo para CD44 (98,92%), CD90 (97,11%) y negativo para el marcador hematopoyético CD45 (0,10%).

Conclusiones: Los condrocitos articulares humanos obtenidos de cinco pacientes ancianos con artrosis mantenían un fenotipo condrocitario y podrían ser potencialmente utilizados para la implantación autóloga. Las condiciones para el cultivo fueron establecidas de acuerdo a los estándares de buenas prácticas de laboratorio para así minimizar el riesgo de contaminación celular in vitro.

Abstract
Objectives: The objective of the present study was to determine if chondrocytes isolated from human cartilage of five elderly patients (middle age 63) with osteoarthritis (stage 3) maintain their proliferation and chondrogenic potential. Isolation and cultivation of chondrocytes was performed according to good laboratory practice (GLP) standards.

Methods: Chondrocytes were isolated from cartilage biopsy by enzymatic digestion. Cultivation of cells was performed in a controlled environment (cleanroom). Phenotype characterization of chondrocytes was achieved by flow cytometry analysis.

Results: Three weeks after cultivation polygonal structures typical for chondrocytes were observed, but spindle/fibroblast like morphology was also detected in culture. Flow cytometric analysis showed that chondrocytes were positive for CD44 (98,35% ± 0,50), CD90 (97,11% ± 0,13) and the cells were negative for hematopoietic marker CD45 (0,21% ± 0,11).

Conclusions: Human articular chondrocytes obtained from five elderly patients with osteoarthritis maintained a chondrocyte phenotype and could be potentially used for autologous implantation. We have standardized the conditions for cultivation according to GLP standards to minimize the risk of in vitro cell contamination.
INTRODUCTION

Cartilage tissue is composed of chondrocytes embedded within a dense extracellular matrix (ECM). It has poor autonomous regeneration capacity, mainly due to its avascular nature. Another factor contributing to poor regenerative capacity of articular cartilage (AC) is the restricted number of ECM particles that can act as repair. The percentage of highly specialized chondrocytes in cartilage tissue is only 1–3% (1). Chondrocytes are unable to migrate to a site of injury; they are able to synthesize fibrous repair tissue, but not sufficiently to fill even small defects (<3 mm in diameter) with a cartilage-like matrix, the defects are not repaired and remains permanently (2). Cartilage was used as one of the first models for research of in vitro engineered tissues and has shown the earliest application for the cell-based therapy mainly due to its cellular homogeneity and avascularity (3).

Chondrocytes within their natural environment actively synthesize and maintain their surrounding matrix. Mature chondrocytes have limited ability to proliferate; they are often mislabeled as dormant. In cell culture, human chondrocytes regain their ability to proliferate. Thus, the basic premise behind autologous Chondrocyte Implantation (ACI) is to overcome the inherent limitations of mature chondrocytes to effectively restore an injured articular surface.

In monolayer culture, these cells respond by undergoing rapid proliferation. Histologically, these chondrocytes reversibly dedifferentiate, assuming a fibroblastic appearance and express type I collagen as opposed to type II collagen normally seen in articular cartilage. Once removed from the monolayer culture and placed in suspension or returned to the articular cartilage environment, the cells undergo a re-differentiation process into normal appearing chondrocytes and again produce type II collagen and proteoglycan aggregates (4).

In 1994 Brittberg et al described the use of ACI in treating full-thickness AC defects of the human knee (5). It was achieved in a two-stage procedure. Stage 1 involved arthroscopic biopsy of healthy AC and cultivation of the chondrocytes to produce between 5 and 10 million cells over a period of 4–6 weeks. Stage 2 involved debridement of the osteochondral lesions and coverage by a periosteal flap followed by open implantation of these cells into the AC defect.

Isolation and cultivation of chondrocytes is a widely spread technique. It is required in ACI as a treatment for osteochondritis dissecans (6), osteoarthritis (7) and articular cartilage injuries produced by trauma (8). In general, it can be used in any articular cartilage injury, where other surgical procedures are not sufficient (4). In vitro chondrocyte manipulation is a crucial phase of autologous chondrocyte implantation. To minimize the risk of in vitro cell contamination, the manipulation must be performed in a controlled environment such as a cleanroom according to good laboratory practice (GLP). The GLP standards provide guidance on implementing GLP requirements critical for laboratory operations (9, 10).

In the current study, we have used basic methods for isolation and cultivation of chondrocytes from human articular cartilage according to GLP standards. Phenotype characterization of chondrocytes was performed by flow cytometry analysis.

MATERIALS AND METHODS

Work in cleanroom according to GLP standards

Isolation and cultivation of chondrocytes was performed in the cleanroom in Associated Tissue Bank of Faculty of Medicine of P. J. Šafárik University and L. Pasteur University Hospital in Košice, Slovakia. A cleanroom is a controlled environment in which the concentration of airborne particles is controlled to specified limits (Tab.1). Contaminants generated by people, process, facilities and equipment were continually removed from the air. Air flow rates, direction, pressurization, temperature, humidity and specialized filtration were all tightly controlled. Persons and materials entered and exited the cleanroom through airlocks (material pass box), gowning rooms and they wore special clothing designed to trap contaminants that are naturally generated by skin and the body. Cells were handled inside biological safety cabinets and cultivated in the gas incubators (Fig. 1).

Isolation and cultivation of chondrocytes

Human cartilage tissue was harvested from the lateral femoral condyle of 5 patients (an average age: 63 years) undergoing total knee replacement surgery due to osteoarthritis (stage 3). Cartilage tissue was harvested in accordance with the ethical standards of L. Pasteur University Hospital committee on human experimentation in Košice, Slovakia. Cartilage tissue was placed into the transport medium containing sterile Autologous Dulbecco’s modified Eagle medium (DMEM; Invitrogen, GIBCO, USA) supplemented with 1% antibiotic/antimycotic solution (10,000 units/mL penicillin, 10,000 µg/mL streptomycin, and 25 µg/mL amphotericin B; Invitrogen, GIBCO, USA) and 1% Insulin-Transferrin-Selenium–A supplement (Invitrogen, GIBCO, USA). Cartilage tissue was minced with a scalpel to small pieces (1x1x1 mm) and digested with 0.1% bacterial collagenase type II (Invitrogen, GIBCO, USA) in Ham’s F-12 (Biochrom AG) for 20 h at 37 °C in 95% air and 5% CO₂ humidified atmosphere. Cell suspension was filtered by 40 µm nylon cell strainer (BD Falcon, Becton, Dickinson and Company) to remove cell raft and matrix debris. The filtrate was then centrifuged at 150 x g for 7 min and the pellet washed twice with DMEM (Invitrogen, GIBCO, USA). Isolated cells were suspended in cell culture medium containing Ham’s F-12 (Biochrom AG), 10% fetal bovine serum (FBS; Invitrogen, GIBCO, USA), 1% antibiotic/antimycotic solution (10,000 units/mL penicillin, 10,000 µg/mL streptomycin, and 25 µg/mL amphotericin B; Invitrogen, GIBCO, USA) and 1% Insulin-Transferrin-Selenium–A supplement (Invitrogen, GIBCO, USA). Chondrocytes were cultivated as a monolayer for expansion in 37 °C humidified incubator with an atmosphere of 95% air and 5% CO₂. The medium was changed 2 times weekly. Confluent layers of chondrocytes were dissociated with 0.05% Trypsin-EDTA solution (Invitrogen, GIBCO®, USA) and the number and viability of cells was assessed by TC10™ Automated Cell Count (Bio-Rad Laboratories).

Characterization of chondrocytes

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Phenotype characterization of chondrocytes after first passage (P1) was performed by flow cytometry. After detaching cells from the tissue culture flasks, the cells were washed twice with phosphate buffered saline solution (PBS; Invitrogen, Gibco, USA) supplemented with 2% FBS (Invitrogen, Gibco, USA). Aliquots of 100,000 cells were incubated with mouse anti-human CD90-PE (Miltenyi Biotec Inc., USA), mouse anti-human CD44-PE (Miltenyi Biotec Inc., USA) and mouse anti-human CD45-FITC (Miltenyi Biotec Inc., USA) for 30 min in the dark. Flow cytometric analysis was performed with FACS Calibur (Becton Dickinson) and CellQuest software (Becton Dickinson). Chondrocytes were considered positive for a surface marker when the percentage of positive cells for that surface marker was ≥ to 95% and cells were considered negative when the percentage of positive cells for that surface marker was ≤ to 5%. The level of marker expression was calculated as the ratio between geometric mean fluorescence intensity of sample cells and that of the negative control.

**RESULTS**

**Chondrocytes isolation and morphology**

Cells isolated from human cartilage after enzymatic digestion were seeded at 75 cm² tissue culture flask (T75) at density 35,000 cells/cm². The morphology of chondrocytes was observed under an inverted phase contrast light microscope (Leica DM IL). Non-adherent or few adherent small round cells were also present in the primary culture after 4 days of cultivation; these cells were removed with first medium change. Heterogeneous population of cells with different morphology was observed after 7 days of cultivation (Fig. 2a). After 21 days of cultivation chondrocytes had more polygonal structure, even though few of cells had spindle/fibroblast like morphology (Fig. 2b). Chondrocytes reached confluence within 3 to 4 weeks. Average number of chondrocytes obtained after trypsinization (P1) was 2,23 x 10⁶ ± 0,5 per mL (Tab. 1).

**Phenotype characterization of chondrocytes**

Flow cytometric analysis showed that chondrocytes were positive for CD44 (98,35% ± 0,50), CD90 (97,15% ± 0,13) after first passage (P1) and the cells were negative for hematopoietic marker CD45 (0,21% ± 0,11) (Fig.3, Tab.1). Data are expressed as means ± SD (n=5).

**DISCUSSION**

The objective of chondrocyte manipulation suitable for autologous implantation is to obtain viable and phenotypically stable cells able to enhance repair processes in the damaged area of human cartilage. Despite the importance of the extracellular matrix, the cells are responsible for the balanced turnover of the extracellular matrix, which is necessary for maintenance of the integrity of the extracellular cartilage. In vitro expansion of chondrocytes is possible via monolayer culture, whereby the cells alter their morphology and metabolism in a process known as dedifferentiation (11). The importance of monolayer expansion of cells from small biopsies is their clinical application in repair strategies such as autologous chondrocyte transplantation (5).

The aim of the present study was to determine if chondrocytes isolated from human cartilage of five elderly patients with osteoarthritis stage 3 maintain their proliferation and chondrogenic potential. It is known that upon digestion of the ECM from a cartilage biopsy and subsequent adhesion to the culturing surface, chondrocytes re-enter the cell cycle and proliferate. Three weeks after cultivation polygonal structures typical for chondrocytes were observed, but spindle/fibroblast like morphology was also detected in culture. Samples derived from aged patients, even in regions of "normal" appearing cartilage, are metabolically not truly normal and

**Table 1. Characterization of chondrocytes isolated from human cartilage of patients with osteoarthritis**

<table>
<thead>
<tr>
<th>Sample</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
<th>Average</th>
<th>SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of cells (x10⁶)</td>
<td>2,53</td>
<td>1,62</td>
<td>2,35</td>
<td>1,81</td>
<td>2,82</td>
<td>2,23</td>
<td>0,50</td>
</tr>
<tr>
<td>Viability (%)</td>
<td>98,00</td>
<td>95,00</td>
<td>99,00</td>
<td>96,00</td>
<td>97,00</td>
<td>97,00</td>
<td>1,58</td>
</tr>
<tr>
<td>CD90 (%)</td>
<td>97,32</td>
<td>97,08</td>
<td>97,11</td>
<td>97,24</td>
<td>97,01</td>
<td>97,15</td>
<td>0,13</td>
</tr>
<tr>
<td>CD44 (%)</td>
<td>98,00</td>
<td>98,13</td>
<td>98,97</td>
<td>97,87</td>
<td>98,80</td>
<td>98,35</td>
<td>0,50</td>
</tr>
<tr>
<td>CD45 (%)</td>
<td>0,12</td>
<td>0,10</td>
<td>0,32</td>
<td>0,30</td>
<td>0,23</td>
<td>0,21</td>
<td>0,10</td>
</tr>
</tbody>
</table>
that chondrocyte phenotype is not stable in vitro, in particular in monolayer culture (12). Most data also suggest that the major phenotypic alterations are initially observed in the superficial zone of early-stage osteoarthritic cartilage, where chondrocytes express de novo abnormal, non-chondrocytic genes; in particular, they express the enzymes required for degrading the matrix that surrounds the cells as well as many of the cytokines and growth factors relevant for turning on the catabolic processes within cartilage (13).

Immunophenotype characterization of chondrocytes was performed by flow cytometry, which offers the possibility to assess and quantify a large number of epitopes on single cells within a short period of time. Immunophenotypic analysis of cells isolated from solid tissues through enzymatic digestions might be compromised due to the reduction or even total loss of surface molecules sensitive to enzymatic treatment. Collagenase type II used in our study did not impair the detection of the CD90, CD44 markers in phenotype of chondrocytes (1P) as was previously reported (14, 15, 16). The hyaluronan receptor CD44 belongs to the polymorphic family of CD44 glycoproteins, which have been implicated in a variety of cellular functions including adhesion to hyaluronan and collagen, and which is present in normal chondrocytes (15). It was previously reported that CD90 is also expressed in a minority of chondrocytes in normal articular cartilage (16). Up-regulation of markers on chondrocytes regarded as distinctive for mesenchymal stem cells (CD90 among others) during monolayer culture suggested that dedifferentiation leads to reversion to a primitive phenotype (17). Hematopoietic marker CD45 known as the leukocyte common antigen was not expressed on chondrocytes. CD 45 should not be present in normal chondrocytes but can appear in dedifferentiated ones (15).

It is important to enhance that cells used in this study were not applied in clinical practice and basic characterization of chondrocytes was the only aim. Nevertheless, weakness of this study is that we did not include a control of healthy articular cartilage and from younger patients with osteoarthritis due to the limitation of donors. In spite of, some investigators reported irreversible changes in phenotype between chondrocytes isolated from OA cartilage versus those of young healthy joints (18, 19, 20). Other reports indicate comparable proliferation or differentiation potential of OA chondrocytes (21, 22).

CONCLUSION

Basics methods of in vitro isolation and cultivation of chondrocytes from OA cartilage were realized in this work. The present study contains also some limitations. More precise characterization of surface and intracellular markers of chondrocytes and comparison with a control group of young healthy patients should be done. In conclusion, we can assume that human articular chondrocytes obtained from elderly patients with osteoarthritis (stage 3) maintained a chondrocyte phenotype and could be potentially used for autologous implantation. We have standardized the conditions for cultivation to minimize the risk of in vitro cell contamination according to GLP standards since chondrocyte manipulation for autologous implantation requires standardised protocols ensuring that the cell product is therapeutically effective and safe.

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REFERENCES


