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Human Cartilage Engineering in an *In Vitro* Repair Model Using Collagen Scaffolds and Mesenchymal Stromal Cells

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Abstract

The purpose of this study was to investigate cartilage repair of *in vitro* lesion models using human bone marrow mesenchymal stromal cells (hBMSCs) with different collagen (Col) scaffolds.

Lesions were made in human cartilage biopsies. Injured samples were pre-treated with interleukin 1 β (IL1 β) for 24 h; also, samples were not pre-treated. hBMSCs were seeded on different types of collagen scaffolds. The resulting constructs were placed into the lesions, and the biopsies were cultured for 2 months in chondrogenic medium.

Using the modified ICRSII scale, neotissues from the different scaffolds showed ICRS II overall assessment scores ranging from 50% (fibrocartilage) to 100% (hyaline cartilage), except for the Col I +Col II +HS constructs (fibrocartilage/hyaline cartilage, 73%). Data showed that hBMSCs cultured only on Col I +Col II +HS scaffolds displayed a chondrocyte-like morphology and cartilage-like matrix close to native cartilage. Furthermore, IL1 β pre-treated biopsies decreased capacity for repair by hBMSCs and decreased levels of chondrogenic phenotype of human cartilage lesions.

Key words: Regenerative Medicine, Tissue Engineering, Hyaline Cartilage, Tissue Scaffolds, Mesenchymal Stromal Cells, Osteoarthritis

Introduction

It is widely accepted that three-dimensional (3D) *in vitro* cultures simulate the *in vivo* situation better than two-dimensional ones. Although micromass culture is considered a useful tool for studies of chondrogenesis, it is not suitable for cell therapy [1]. Scaffolds can, however, provide an appropriate 3D environment for cell viability and proliferation, cell differentiation and maintenance of a specific phenotype [2].

Tissue Engineering using biodegradable scaffolds, cells and cell factors has evolved into a multidisciplinary field with an aim to develop biological substitutes with biochemical, structural, morphological and functional properties similar to native cartilage for subsequent use for *in vivo* treatment [3]. The most important issue in Tissue Engineering is the selection of cell type and optimal scaffold.

Most studies have tested growth and differentiation of cells on various scaffolds but have not placed the constructs inside a native cartilage environment or taken the host tissue into account [4]. *In vitro* testing is easier than *in vivo* testing for obtaining standardized and quantifiable information about cell cytotoxicity, proliferation and differentiation capacity [5]. This model allows us to

analyze human cartilage samples from a single donor using different study variables and to test different constructs in an *in vitro* native environment. In our study, this model was used to test the repair capacity of different constructs.

Materials and Methods

Isolation and culture of bone marrow stromal cells

Each donor in the study gave informed consent according to the guidelines of the local ethics committee and principles of Declaration of Helsinki. This study was approved by the Ethics Committee of Clinical Investigation of Galicia (Santiago de Compostela, Spain).

Bone marrow cells were obtained from three hip fracture patients who underwent total hip replacement, as previously described [6]. Cells were extracted by washing the bone marrow with culture medium: Dulbecco's Modified Eagle's Medium (DMEM; Lonza, Spain) with 20% foetal bovine serum (LabClinics, Spain). The isolated cells were cultured in a 5% humidified CO₂ atmosphere at 37°C. The resultant cell suspension was cultured and expanded. We previously confirmed by characterization [6] that cells obtained by this method are hBMSCs.

Scaffolds

Different Col scaffolds (Opocrin S.p.A., Modena, Italy) had been previously tested by our group [6, 7]. From those, scaffolds producing better chondrogenic hBMSC phenotypes, including type I collagen (Col I), Col I and heparan sulfate (Col I+HS), Col I and type II collagen (Col II) and HS (Col I+Col II+HS) and Col I and Col II and chondroitin sulfate (Col I+Col II+CHS) were selected. The hBMSCs were cultured on the surface of 4 mm-diameter sponges and incubated for 1 h at 37°C. These constructs were used for the *in vitro* lesion model using chondrogenic medium: hMSC Chondrogenic Differentiation Bulletkit medium (Lonza, Spain) with 10ng/ml of human transforming growth factor β -3 (TGF β -3, Prospec-Tany Technogene Ltd., Israel).

In vitro lesion model

Cartilage samples were obtained from three patients who underwent total hip or knee replacement because of fracture or osteoarthritis (OA). The cartilage was sliced with a scalpel and 6mm biopsies were obtained using disposable biopsy punches (Kai Medical, Germany).

3 mm-Diameter lesions were made in human cartilage biopsies using a dental drill (Gebr. Brasseler Gmbh & Co. KQ, Germany) with a rotor (EWL K9, Germany). Over a two-month period, two experiments were performed:

1) Non-IL1 β Group: 2x10⁵ cells were seeded on scaffolds. These constructs were placed into the lesion and chondrogenic medium was added. The medium was changed every 3-4 days.

2) IL1 β Group: Cartilage biopsies with lesions were pre-treated with 10ng/ml of IL1 β (Sigma, USA) for 24 hours. 2x10⁵ cells were seeded on scaffolds, the resulting sponge-constructs were introduced into the lesion and chondrogenic medium was added. The medium was changed every 3-4 days.

Following controls were included: defect without constructs, defect with scaffolds and without cells, and native cartilage without defect (Figure S1).

Histological analysis

The resulting neotissue was evaluated histochemically using 4 µm-thick deparaffinized sections. These sections were then stained with hematoxylin and eosin (HE), and stained for collagen and proteoglycans (PGs) with Masson's trichrome (MT) or Safranin O (SO), respectively. The stained sections were examined using an optical microscope (Olympus BX61, USA) equipped with a digital camera (Olympus DP70, USA). Quantitative analyses of the SO staining were measured using ImageJ 1.48v (*National Institutes of Health, Bethesda, USA*). After the colour substraction of non-stained areas, the percentage of metachromatic areas was measured and expressed as mean±standard error.

To assess the quality of cartilage repair, the ICRS II histological scoring system [8] was used with some modifications to adapt it for analysis of in vitro cartilage repair. The neotissues formed were scored by three blind observers. Our modified score system comprised 7 out of the 14 parameters: "cell morphology" and "chondrocyte clustering" as well as "surface architecture" and the "basal integration" of the neotissue formed were observed by the HE staining; "tissue morphology" was assessed by analyzing the collagen fiber distribution with MT staining; the parameter to evaluate PG content was "matrix staining", which was assessed using SO staining; and, the "overall assessment" was the statistical mean of the values for all 6 parameters analyzed.

Further immunohistochemical analysis of Col I, Col II and aggrecan (Agg) was performed in samples with highest ICRS II score.

Results

Controls and Non-IL1 β model

Control constructs without cells did not show

presence of neotissue formation and the scaffolds were not degraded (Figure S1, Ctrl w/o cells). Self-repair did not occur in the negative control (Figure S1, control lesion). In the Non-IL1 β model, histochemistry after two months revealed that cells seeded on all scaffolds filled almost 100% of the lesion and that neotissues were integrated with the border of the lesion (Figure 1, H-E Non-IL). Scaffolds were almost fully degraded in this model, except the Col I+HS scaffold, and cellular density in all the grafts was higher (Figure 1, H-E Non-IL) than that in the native cartilage (Figure S1, positive control). Spindleand round-shaped cell morphology was found in all neotissues.

Staining with MT showed the presence of Col in the ECM of all the constructs (Figure 1, H-E Non-IL), that was observably more homogeneous in the tissues of the Col I+Col II+HS constructs. The presence of PGs detected by SO staining (Figure 1, H-E Non-IL) was more metachromatic in the Col I+Col II+HS constructs (52.426±4.877) than in Col I (5.107±3.337), Col I+HS (23.406±6.189) and Col I+Col II+CHS (32.722±8.781) constructs.



Figure 1. Histology of repair in human cartilage non interleukin- 1β-treated lesions (Non-IL) and IL1β-treated lesions (IL). Hematoxylin-eosin (HE), Safranin O (SO) and Masson's Thricrome (MT) staining of the models. Scale bar 100µm (smaller images 200 µm). Col I: type I collagen; Col II: type II collagen; HS: heparan sulfate; CHS: chondroitin sulfate.



Figure 2. Histology and immunohistochemistry of repair obtained using type I and II collagen and heparan sulfate scaffolds (Col I+Col II+HS), in non-interleukin-1β-treated lesions. Images of Safranin O (SO) and Masson's Thricrome (MT) staining, and type I (Col I) and II (Col II) collagen and aggrecan (Agg) immunostaining performed on the neotissues formed on Col I+Col II+HS scaffolds in the Non-IL model. Col and Agg immunostaining were counterstaining with hematoxylin-eosin. Scale bar 100µm.

Table 1. Assessment of quality of repair in human cartilage non interleukin- 1β -treated lesions (Non-IL) and IL1 β -treated lesions (IL). International Cartilage Repair Society II (ICRS II) scale values for different scaffolds in the Non-IL1 β -treated and IL1 β -treated model. Our modified scoring system comprises 7 of the 14 original parameters.

Parameter	Col I (Non-IL)	Col I (IL)	Col I +HS (Non-IL)	Col I +HS (IL)	Col I + Col II +HS (Non-IL)	Col I + Col II +HS (IL)	Col I +Col II +CHS (Non-IL)	Col I +Col II +CHS (IL)
Tissue morphology	30	5	35	30	50	20	35	20
Matrix staining	0	0	5	0	50	5	15	0
Cell morphology	15	0	5	25	70	40	80	40
Chondrocyte clustering	100	100	100	100	100	100	100	100
Surface architecture	90	80	80	90	90	90	40	50
Integration	30	5	60	50	80	65	10	50
Overall assessment	44%	31%	47%	49%	73%	53%	46%	43%

Col I: type I collagen; Col II: type II collagen; HS: heparan sulfate; CHS: chondroitin sulfat.

By macroscopic ICRS II assessment of the Non-IL model, the repair score was between 44% and 47%, or fibrocartilage, for Col I, Col I+Col II+CHS and Col I+HS scaffolds. Col I+Col II+HS scaffolds showed a chondrogenic phenotype, 73%, or mixed firbrocartilage and hyaline cartilage on the ICRS II scale. This was confirmed by histological and immunohistochemical analysis that showed the presence of PGs and Col, being positive for Col II and Agg (Figure 2).

IL1β-treated model

In the IL1 β model, HE staining showed neotissue formation within the lesion with fewer rounded cells than in the Non-IL1 β model (Figure 1, H-E IL). The presence of PGs detected by SO staining (Figure 1, H-E IL) was more metachromatic in the Col I+Col II+CHS constructs (23.228±1.704) than in Col I (5.738±1.283), Col I+HS (5.107±2.435) and Col I+Col II+HS (20.973±6.849) constructs. These morphological changes with the loss of ECM components observed by SO and MT staining resulted in lowering of the ICRS II scores except those for the Col I+HS and Col I+Col II+CHS constructs (49%-43%, respectively) (Table 1).

Discussion

This model supposes an improvement compared with other 3D *in vitro* models.

Because cell phenotype and extracellular matrix (ECM) vary with cartilage depth [9], constructs, once implanted, are influenced by the native cell metabolism of each zone [10].

We found that cellular density in all the grafts was higher than in the native cartilage and, flattened-like undifferentiated cells were observed in all the scaffolds. The presence of undifferentiated hBMSCs or chondrocyte-like cells without zonal organization may be advantageous for Cartilage Engineering because immature tissue is capable of maturing when implanted in the joint [4].

Col I+Col II+HS scaffolds showed a fibrocartilage/hyaline cartilage phenotype, using the ICRS II scale. These results are similar to those obtained in a clinical study following microfracture treatment [11].

Chondrocytes implanted into focal or OA joint lesions were susceptible to native inflammation. Furthermore, problems with mechanics in OA and eventual loss of integrity, functionality and, long-term neotissue viability could be compromised [12]. Khan *et al.* found that a single short catabolic pulse of IL1 β followed by post-culture anabolic conditions is sufficient to generate mechanically robust, integrative cartilage repair [13]. That is why secondly, an *in vitro* lesion model in which explants were pre-treated with IL1 β before culture in chondrogenic medium was developed.

Different studies have shown that *in vivo* infiltration of IL1 β into the joints results in degradation of Col and PGs from the chondrocyte ECM [13]. In our study, constructs showed lower amounts of PGs and less rounded cells after IL1 β treatment than without treatment, except in CHS scaffolds. Scaffolds composed of CHS have been described as having better anti-inflammatory properties [14].

Regarding to the ICRS II scale, it was demonstrated lower inter- and intra-reader variability compared with other traditional scales. The overall assessment and matrix staining scores had the best correlation coefficients for inter- and intra-reader variability [8]. However, there are limitations of this study to take into account. It is necessary to consider any disadvantage of assessing the quality of repair in the *in vitro* developed models using the ICRS II scale, which was originally designed to evaluate in vivo osteochondral repair. ICRS II [8] was developed comprising 14 criteria: tissue morphology, matrix staining, cell morphology, chondrocyte clustering, surface architecture, basal integration, tidemark formation, subchondral bone fibrosis, inflammation, calcification, vascularization, abnormal surface assessment, mid/deep zone assessment and overall assessment. However, we focus on the assessment of 7 out of 14, which were related to chondrocyte phenotype and tissue structure. The assessment of the other 7 parameters was not possible in our developed model (tidemark formation, subchondral bone fibrosis, inflammation, abnormal calcification, vascularization, surface assessment and mid/deep zone assessment).

It is also relevant to point out that lack of mechanical stimuli negatively influences the quality of the neotissue formed. It is well known that continuous passive movement was shown to stimulate the repair of focal lesions with a neotissue similar to hyaline cartilage [9] and movement and loading in the joint were demonstrated to be essential for development and maintenance of normal cartilage structure [15]. The duration of our study was only two months, which is less time than that described in the literature. Because cartilage repair is a slow process, mathematical models have predicted that more than 18 months would be required to obtain complete cartilage maturation *in vivo* [5]. *In vitro* culture for long

periods of time increases the risk of culture contamination and cartilage degradation [3]. However, our group had previously tested these two-month-culture-models without culture contamination or tissue degradation [16].

In conclusion, in an *in vitro* model using hBMSCs cultured on a Col I +Col II +HS scaffold displayed higher overall human cartilage lesion repair assessment than those of the other scaffolds. The capacity for repair by hBMSCs and the levels of construct chondrogenic phenotype of human cartilage lesions are lessened by pretreatment with IL1 β .

Supplementary Material

Figure S1. http://www.medsci.org/v14p1257s1.pdf

Abbreviations

hBMSCs: human bone marrow mesenchymal stromal cells; Col: collagen; IL1_β: interleukin 1_β; Col I: type I collagen scaffolds; Col I+HS: Col I and heparan sulfate scaffolds; Col I+Col II+HS: Col I, Col II and HS scaffolds; Col I+Col II+CHS: Col I, Col II sulfate and chondroitin scaffolds; TGF β -3: transforming growth factor β -3; ICRS: International Cartilage Repair Society; 3D: three dimensional; OA: osteoarthritis; DMEM: medium Dulbecco's Modified Eagle's Medium; ECM: extracellular matrix; HE: hematoxylin and eosin; MT: Masson's trichrome; SO: Safranin O; PGs: proteoglycans; Agg: aggrecan.

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Competing Interests

The authors have declared that no competing interest exists.

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