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Effects of Dynamic Fluid Pressure on Chondrocytes Cultured in Biodegradable Poly(glycolic acid) Fibrous Scaffolds

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INTRODUCTION

Cartilage defects cause considerable pain for the individual patient and they inflict a heavy burden on society. An estimated 70 million Americans are affected by arthritis and related conditions and the total cost for treating these diseases is $82 billion annually. The clinical problems range from small traumatic osteochondral defects to global destruction of the joint surface from arthritis. Damage to articular cartilage often requires surgical intervention because of its limited capacity for self-repair. The unique challenges of each clinical situation call for different treatment strategies. The current treatment options include shaving the articular surface, perforating the underlying subchondral bone, periosteal and perichondral grafts, chondrocyte transplantation, osteotomy, and artificial joint replacement. However, none of these techniques can effectively manage the symptoms and predictably restore normal cartilage function.

Tissue-engineering strategies using biodegradable polymers as scaffolds for cell transplantation provide many advantages over current therapies. The three components common to many cartilage tissue-engineering approaches are cells (either chondrocytes or chondroprogenitor cells), a scaffold, and various soluble and extracellular matrix signals. In addition to serving as a substrate, the scaffold can interact with the other elements in a dynamic and synergistic manner to direct and organize the process of regeneration.

Various synthetic materials have been explored to serve as scaffolds for cartilage tissue engineering. Injectable formulations based on poly(ethylene oxide), poly(anhydride), and oligo-poly(ethylene glycol)-fumarate may be delivered through minimally invasive procedures to treat contained focal defects. Synthetic polymers as preformed scaffolds may be preferable for replacing large, noncontained defects. Among them, highly porous poly(glycolic acid) (PGA) nonwoven fibrous scaffolds have been used extensively for chondrocyte culture and subsequent transplantation. These studies have shown that PGA provides a suitable substrate for chondrocyte proliferation and cartilage tissue formation both in vitro and in vivo. However, the quality of the cartilage tissue formed in synthetic scaffolds is inferior to the natural cartilage in both structural organization and biomechanical properties.

The role of mechanical force in regulating chondrocyte function and maintaining normal cartilage tissue has long been recognized. During normal joint movement, cyclic forces generate complex and dynamic stress and strain fields throughout articular cartilage. At present, continuous passive motion (CPM) is used clinically in postoperative care after cartilage repair. CPM transmits forces to cartilage through direct deformation at the opposing cartilage surfaces in the knee joint as well as through normal and tangential fluid-induced forces. The focus of this study was on the role of hydrostatic force, the normal component of the fluid-induced forces experienced during CPM, on chondrocytes cultured in biodegradable polymer scaffolds. Specifically, we investigated the effects of dynamic fluid pressure (at a magnitude of 13 kPa and a frequency of 0.3 Hz) and the time course of pressurization on cultured chondrocyte–PGA constructs during a 6-week period. Low-level oscillations in hydrostatic pressure (13 kPa at 0.3 Hz) were chosen because they are similar to the cyclic oscillations of intraarticular synovial fluid pressure during CPM and because cultured chondrocytes were shown to respond to such a mechanical stimulus.
MATERIALS AND METHODS

Articular cartilage harvest and chondrocyte isolation

Articular cartilage was harvested aseptically from the femoropatellar grooves of 2-month-old New Zealand White rabbits. This protocol was approved by the Institutional Animal Care and Use Committee at the Mayo Clinic (Rochester, MN). Chondrocytes were isolated by digestion with a collagenase II solution (Worthington Biochemical, Lakewood, NJ). The primary cells were expanded in medium consisting of Dulbecco’s modified Eagle’s medium (DMEM), 10% fetal bovine serum, 2 mM L-glutamine, 0.1 mM nonessential amino acids, L-proline (50 μg/mL), 1 mM sodium pyruvate, and gentamicin (25 μg/mL) (all from Invitrogen GIBCO, Grand Island, NY), as previously described.

Chondrocyte seeding and culture in polymer scaffolds

Poly(glycolic acid) (PGA) nonwoven fibrous disks were purchased from Albany International Research (Mansfield, MA). These disks were 5 mm in diameter and 2 mm thick, with a fiber diameter of 13 μm, a void volume of 97%, and a bulk density of 62 mg/mL. Before use in cell culture, PGA disks were sterilized with ethylene oxide and prewetted by sequential immersion in 70% ethanol, phosphate-buffered saline (PBS; Invitrogen GIBCO), and chondrocyte culture medium (see Articular Cartilage Harvest and Chondrocyte Isolation, above).

Chondrocytes at first passage were seeded at a density of 5 × 10^6 cells per scaffold in the prewetted PGA disks placed in 12-well plates (non-cell culture treated; Fisher, Pittsburgh, PA) according to the method described by Schreiber et al. The cell–polymer constructs were cultured for a total of 6 weeks with the culture medium changed every 2 to 3 days. Insulin and ascorbic acid at 50 μg/mL were added at each feeding.

The chondrocyte–PGA constructs were divided into four sample groups, identified by culture condition (S, static; D, dynamic fluid pressure) and the number of weeks for each condition. After 1 day of cell seeding, the constructs were cultured for 6 weeks under static conditions (Static), 1 week of static culture and then 5 weeks under dynamic fluid pressure (S1/D5), 2 weeks of static culture and then 4 weeks under dynamic fluid pressure (S2/D4), or 6 weeks under dynamic fluid pressure (DFP). During the fluid pressure culture period, samples were pressurized at a magnitude of 13 kPa and a frequency of 0.3 Hz for a total of 4 h/day. All constructs were placed in the pressure chamber for 4 h/day and all cells were allowed to experience the purge cycle.

Application of dynamic fluid pressure

Dynamic fluid pressure was applied to the chondrocyte–PGA constructs by compression of the gas phase above the culture medium covering the constructs, using a custom-built apparatus similar to that previously described. As shown schematically in Fig. 1, tissue culture plates containing the cell–polymer constructs in culture medium were placed in a rigid, cylindrical stainless steel pressure chamber. A pneumatically driven membrane chamber was used to create dynamic pressurization in the gas phase in the attached pressure chamber. Pressure transducers were placed in the gas phase to measure the actual pressure in the chamber. A digital data acquisition and analysis system with a sample frequency of 150 Hz was used to monitor the response. A 5% CO2 level in the gas phase was maintained in the pressure chamber. Using this system, a cyclic, squarewave dynamic fluid pressure at a magnitude of 13 kPa and a frequency of 0.3 Hz was applied to the constructs for 4 h/day.

Sample analysis

After 3 weeks and 6 weeks of culture, five chondrocyte–PGA constructs from each sample group (Static, S1/D5, S2/D4, and DFP) were harvested: three specimens for biochemical analysis, one for histologic evaluation, and one for scanning electron microscopy (SEM).

Biochemical analysis. The diameter and wet weight of the cell–polymer constructs were measured. The construct dry weight was obtained after drying with a SpeedVac (model SC200; Thermo Electron, Waltham, MA). A portion of the dried constructs was reserved for the collagenotyping assay, whereas the rest was digested with a papain solution (Sigma, St. Louis, MO) as previously described for further biochemical analysis. Total DNA in the constructs was measured fluorometrically with a Hoechst 33258 (Polysciences, Warrington, PA) dye-binding assay. A conversion factor of 7.7 pg of DNA per chondrocyte was used to obtain the total cell number.

FIG. 1. Schematic of the dynamic fluid pressure system.
Total collagen in the constructs was determined from the measured hydroxyproline content after acid hydrolysis, using a hydroxyproline-to-collagen ratio of 1:20. The percentage of type II collagen in the total collagen was quantified by comparing the spectrophotometric densities of specific cyanogen bromide peptide bands on sodium dodecyl sulfate–polyacrylamide gels with a standard curve. Glycosaminoglycan (GAG) content in the cellular constructs was determined by the dimethylene blue dye-binding assay.

**Histology.** For histologic evaluation, the constructs were fixed in 10% neutral buffered formalin (Sigma), dehydrated, and embedded in methyl methacrylate. Tissue blocks were sectioned with a microtome into 8-μm sections and stained with safranin O for GAG distribution.

To quantify the relative amount of cartilage tissue present in the cell–polymer constructs, at least five sections obtained from various depths throughout the construct were analyzed with OsteoMeasure software (Osteometrics, Decatur, GA). The percentage of cartilage matrix was defined as the surface area stained red divided by the total surface area.

**Scanning electron microscopy.** The three-dimensional morphology of the cell–polymer constructs after 6 weeks of culture was assessed by SEM. On harvest, the constructs were washed with PBS, fixed in 2.5% glutaraldehyde (Sigma), and stained with 1% osmium tetroxide (Polysciences). The stained cell samples were dried in a critical point dryer, mounted on metal stubs with double-sided carbon tapes, and coated with a 50:50 mixture of gold and platinum. Both construct surfaces and freeze-fractured cross-sections were examined with a Hitachi S4700 field emission scanning microscope (Hitachi High Technologies America, Pleasanton, CA).

**Statistical analysis.** All data are reported as means ± standard deviations (SD) for n = 3 experiments, except for histologic quantification, where at least five sections were measured. Single-factor analysis of variance (ANOVA) was used to assess the statistical significance of results. The Schefé method was employed for multiple comparison tests at significance levels of 95%.

**RESULTS**

**Construct size, dry weight, and water content**

The size and weight of the chondrocyte–PGA constructs after 3 and 6 weeks of culture under Static, S1/D5, S2/D4, or DFP conditions were measured. The diameter of constructs was found to increase from 5.35 ± 0.12 mm for initial PGA disks to 6.73 ± 0.27 mm after 6 weeks, indicating outward tissue growth from the scaffolds.

The dry weight of all cell–polymer constructs after incubation for 3 weeks was 4.33 ± 0.25 mg, more than twice the initial weight of the scaffold (1.70 ± 0.27 mg), and nearly twice the weight of the non-cell-seeded polymer scaffolds incubated in culture medium (2.23 ± 0.42 mg). The slight increase was probably due to absorption by the scaffold of serum proteins in the culture medium. By 6 weeks in culture, the construct dry weight for all sample groups had increased to 6.34 ± 0.16 mg. In contrast, PGA scaffolds without cells had completely disintegrated after 6 weeks and could not be retrieved for further evaluation.

The water content of the constructs, calculated from their wet and dry weights, was 88.7 ± 1.5% for all sample groups after 3 or 6 weeks of culture. The water content of the constructs was in the range reported for native cartilage.

**Cell density**

Cell number in the chondrocyte–PGA constructs was determined by the DNA assay and normalized to the construct dry weight (Fig. 2). All sample groups had similar (p > 0.05) cell densities at the two time points tested. By 6 weeks, the cell densities had increased to 106.7, 120.0, 108.1, and 96.8 (×10⁷ per gram of construct dry weight) for static, S1/D5, S2/D4, and DFP constructs, respectively. No significant (p > 0.05) increase in cell number from 3 to 6 weeks was found, except for the S1/D5 samples.

**Collagen**

Total collagen in the cell–polymer constructs was determined by hydroxyproline assay. Of the four sample groups studied, the collagen content as a percentage of construct dry weight was less than 10% after 3 weeks, but increased 2-fold after 6 weeks (Fig. 3A). Total collagen at 6 weeks (23.7, 20.2, 19.0, and 19.9% for the Static, S1/D5, S2/D4, and DFP groups, respectively) was about one-third that of native cartilage. Between 90 and 95% of the total collagen was type II collagen, as determined by collagen-typing assay (Fig. 3B). No significant (p > 0.05) difference between sample groups was observed. The percentage of collagen type II in the constructs was similar to that for normal cartilage.

**GAG**

Glycosaminoglycan content as a percentage of construct dry weight was determined by dimethylene blue assay. As shown in Fig. 4, the GAG contents in the four sample groups were similar, increasing significantly (p < 0.05) from approximately 8% at 3 weeks to 18% after 6 weeks. The values at 6 weeks were approximately half that for native cartilage.
Histology

Chondrocyte–PGA constructs cultured under Static, S1/D5, S2/D4, and DFP conditions were stained with safranin O for GAG distribution after 6 weeks. The periphery and middle portions of the constructs are shown, respectively, in the top and bottom panels of Fig. 5. Intense red staining at construct periphery indicated the production of a sulfated cartilage matrix for all sample groups (Fig. 5, top). The chondrocytes had a rounded morphology and resided inside lacunae. At the center of the constructs, however, staining was less intensive and polymer fibers not yet degraded were still present (Fig. 5, bottom).

The relative amount of cartilage matrix in the tissue sections was quantified by normalizing the red-stained area to the whole section area. The results showed that between 94.8 and 97.9% of all cell–polymer constructs was cartilage matrix after 6 weeks.

Morphology

The three-dimensional morphology of the tissue formed in the PGA scaffolds was similar to that of native cartilage after 6 weeks. Figure 6 shows representative images of chondrocytes cultured for 2 weeks under static conditions followed by 4 weeks under DFP conditions (S2/D4 group). Chondrocytes were seen to cover the tissue surface (Fig. 6a and b); and the cells were spherical in shape and embedded in their extracellular matrix. A higher magnification view of the cross-sections shows a chondrocyte residing in its lacuna and surrounded by a highly fibrous extracellular matrix (Fig. 6c). A remaining PGA fiber was also seen adjacent to the cartilage matrix (Fig. 6d).

DISCUSSION

This study was designed to investigate the effects of dynamic fluid pressure on chondrocytes cultured in biodegradable PGA scaffolds. We investigated the effects of pressure application (at a magnitude of 13 kPa and a frequency of 0.3 Hz) as well as the time course of pressurization (4, 5, or 6 weeks with 4 h of pressurization per day).
Cartilage matrices were developed in the chondrocyte–PGA constructs under all conditions studied. The water content in these constructs was about 90%, similar to native cartilage. The dry constructs were composed of approximately 18% GAG and 20% collagen after 6 weeks of cultivation. Chondrocytes and remaining polymer fibers made up the majority of the rest of the construct dry weight. We showed that no significant difference in chondrocyte proliferation and cartilage matrix formation in the cell–polymer constructs was found between sample groups subjected to static culture (Static) or various time periods of dynamic fluid pressurization (S2/D4, S1/D5, and DFP).

The initial cell-seeding density used in this study was $5 \times 10^6$ cells per scaffold, corresponding to $300 \times 10^7$ cells/g of dry scaffold. Intermittent pressurization did not affect cell numbers compared with the static control (Fig. 2). Freed et al.$^{26}$ seeded bovine calf articular chondrocytes in similar PGA fibrous disks at the same initial cell density. They observed that the construct cellularity reached a plateau within 6 days of culture in a rotating bioreactor, reaching $15 \times 10^6$ cells per construct. Our results also showed that cellularity leveled off between 3 and 6 weeks of culture for most sample groups. Carver and Heath$^{27}$ used similar fibrous PGA mesh to seed foal articular chondrocytes and subjected the constructs to an intermittent hydrostatic pressure of 3.4 MPa, 250 times higher than the pressure used in our study. The cell numbers increased with culture times of 3 to 5 weeks but was not affected by pressurization. Therefore, our results confirm previous findings that cell numbers are not significantly affected by intermittent pressure. The cell number in our study after 6 weeks of culture with or without pressurization ranged between $10 \times 10^7$ and $12 \times 10^7$ cells/g wet weight, close to the native chondrocyte cell number of $4 \times 10^7$ to $6 \times 10^7$ cells/g wet weight.$^{27}$

The total collagen percentage based on construct dry weight as a function of culture time is shown in Fig. 3A. No significant differences in collagen content were observed between the static and pressurized groups; however, the total collagen content increased by 3- and 2.7-fold for these groups, respectively, with increasing culture time from 3 to 6 weeks. It was also observed that total collagen content decreased slightly from 24% for static cultures to 20% for dynamic cultures. In previous studies, Carver and Heath$^{27}$ found a 1.5-fold increase in total collagen content.
for foal chondrocytes after 5 weeks of culture with or without pressurization at 3.4 MPa. Moreover, they observed a slight decrease in total collagen for all time points when constructs were pressurized at 3.4 MPa, which is consistent with our findings with intermittent pressurization of 13 kPa. In another experiment, enhanced collagen synthesis by equine chondrocytes in PGA scaffolds was found under an intermittent pressure of 6.8 MPa but not 3.4 MPa, suggesting that a high pressure level may be needed to stimulate collagen formation.

The percentage of type II collagen was higher than 90% for static as well as dynamic constructs after 3 and 6 weeks of culture (Fig. 3B). There was a slight (statistically insignificant) decrease in percent collagen type II from 95 to 91% when intermittent pressure was applied to the constructs. The finding of a high type II collagen content demonstrates that the chondrocyte phenotype was maintained in the constructs in the presence of a dynamic fluid pressure of 13 kPa. The percent type II collagen for dynamic constructs (91%) was comparable to previously reported levels for chondrogenesis in a PGA bioreactor system in the absence of pressure and to the level for natural bovine articular cartilage (90.3%). In another study, by Ikenoue et al. intermittent hydrostatic pressure was applied to normal human articular cartilage with pressure levels of 1, 5, and 10 MPa for 4 h/day for either 1 day (4 × 1) or 4 days (4 × 4). They observed no change in expression pattern for collagen II for 4 × 1 loading regardless of the pressure level. However, the collagen II expression level was upregulated with 4 × 4 regimens only for pressure levels of 5 and 10 MPa. Therefore, consistent with our findings, collagen II production is not influenced by a short duration or low magnitude of intermittent hydrostatic pressure.

The GAG content in the constructs approximately doubled as the culture time was increased from 3 to 6 weeks (Fig. 4). A slight (statistically insignificant) decrease in GAG content from 21% of dry construct for static cultures to 16–18% of dry construct for dynamic cultures was observed. The GAG content after 6 weeks of culture with intermittent pressure was 18% of dry mass, which was significantly less than that of native articular cartilage (41%). Carver and Heath found that an intermittent pressure of 3.4 MPa significantly increased GAG content, exhibiting native GAG levels after 6 weeks.

As pointed out in introduction, we chose a pressure magnitude of 13 kPa and a pressure frequency of 0.3 Hz to mimic the normal component of the fluid pressure during continuous passive motion. This pressure regimen has been shown previously to stimulate cellular activity in cultured periosteal cells. However, our results indicate that an intermittent hydrostatic pressure of 13 kPa has no
effect on cellularity, total collagen, and GAG content of the chondrocytes cultured in PGA scaffolds. Analysis of joint loading profiles shows that mechanical forces that produce either dilatational or deviatoric stresses exhibit differential effects on articular chondrocyte metabolism. The loading mode in this study was normal hydrostatic pressure with no direct deformation or fluid flow or deformation-induced shear stresses transmitted to the seeded chondrocytes. Future experiments are required to investigate the effects of direct deformation and fluid shear, in order to further elucidate the effects of continuous passive motion on chondrogenesis of cell–PGA constructs.

CONCLUSIONS

We have shown that cartilage matrices were produced in chondrocyte–PGA constructs after 6 weeks of culture. The amount of type II collagen and glycosaminoglycan in the dry constructs increased significantly from 3 to 6 weeks. The formed matrices were similar to normal cartilage in terms of water content and percentage of type II collagen. We also found that the normal component of the hydrostatic force did not affect cartilage matrix formation under the conditions tested. To further elucidate the role of continuous passive motion, more experiments are needed to study the other forces transmitted to cartilage, such as direct deformation and fluid shear, or the combined effects of dynamic fluid pressure with these forces.

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