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Effects of both vitamin C and mechanical stimulation on improving the mechanical characteristics of regenerated cartilage

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ABSTRACT

The present work describes the influence of both vitamin C (VC) and mechanical stimulation on development of the extracellular matrix (ECM) and improvement in mechanical properties of a chondrocyte–agarose construct in a regenerating tissue disease model of hyaline cartilage. We used primary bovine chondrocytes and two types of VC, ascorbic acid (AsA) as an acidic form and ascorbic acid 2-phosphate (A2P) as a non-acidic form, and applied uniaxial compressive strain to the tissue model using a purpose-built bioreactor. When added to the medium in free-swelling culture conditions, A2P downregulated development of ECM and suppressed improvement of the tangent modulus more than AsA. By contrast, application of mechanical stimulation to the construct both increased the tangent modulus more than the free-swelling group containing A2P and enhanced the ECM network of inner tissue to levels nearly as high as the free-swelling group containing AsA. Thus, mechanical stimulation and strain appears to enhance the supply of nutrients and improve the synthesis of ECM via mechanotransduction pathways of chondrocytes. Therefore, we suggest that mechanical stimulation is necessary for homogeneous development of ECM in a cell-associated construct with a view to implantation of a large-sized articular cartilage defect.

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1. Introduction

Articular cartilage covers the surface of the ends of bones in synovial capsules and performs the important function of high performance weight-bearing at very low friction and wear in daily activities during a healthy life. The behavior of articular cartilage tissue as a mechanical system is dependent chiefly on an extracellular matrix (ECM), which consists primarily of a protein, called type II collagen (10–20% of the wet weight), and proteoglycans (about 10% of the wet weight) [1]. Steric and electrostatic interactions of ECM molecules in the cartilage tissue occur between the cationic collagen fibers and the anionic proteoglycans to provide a highly charged environment under neutral pH conditions. Although adult articular cartilage is a remarkable load-bearing system, following traumatic injury or under conditions of wear and tear, it lacks the ability of self-repair, which can often lead to osteoarthritis (OA).

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Such diseases of hyaline cartilage are a major health problem, especially in industrialized countries with relatively long active life expectancies. At present, no cell-assisted tissue regeneration therapy for the reliable and durable replacement of damaged articular cartilage has been established [2–4]. However, in recent years, tissue engineering has become a promising option for the treatment of OA and has allowed researchers to produce functional replacements for diseased cartilage [5,6]. Developments in therapeutic strategies for damaged cartilage treatment have increasingly focused on the promising technology of cell-assisted repair, which proposes the use of autologous chondrocytes or other cell types to regenerate articular cartilage [4,7].

There is still too little knowledge available to establish a suitable design strategy for reconstructing tissue-engineered cartilage that matches the mechanical properties of natural tissue. In this study, we focused on the relationship between the collagen network and the mechanical properties of a regenerated–cartilage tissue model and showed that the collagen network interconnecting chondrocytes improved the mechanical properties of the tissue to be equivalent to that of scaffold material without chondrocytes if there was no linkage between cells by the ECM network [8]. We used ascorbic acid (AsA), a type of vitamin C (VC), in the culture medium to control collagen synthesis in order to investigate the

effect of developing a collagen network on the mechanical properties of the tissue model. However, high concentrations of AsA were found to be toxic to the chondrocytes with no concomitant improvement in mechanical properties.

As for AsA above, we used ascorbic acid 2-phosphate (A2P), a non-acidic form, for suppressing the cytotoxicity and effective development of the mechanical characteristics of the tissue, and demonstrated the relationship between VCs in the culture medium and development of ECM and mechanical properties of the construct. In addition, we thought that applying a compressive deformation to the tissue model would not only improve diffusion of culture medium into the tissue and accelerate the supply of nutrients, but also activate cell-signaling via mechanosensor and mechanotransduction pathways. It is well known that loading mechanical stimuli to regenerated-cartilage tissue and chondrocytes is distinctly important for both enhancing synthesis of ECM and inducing chondrogenesis differentiation of stem cells [9]. Hence, in order to apply a compressive deformation to the tissue model for development of ECM, we made a purpose-built bioreactor capable of applying an uniaxial unconfined compressive strain to individual constructs in an incubator. The purpose of the present study was to investigate the effects of applying both two types of VC and dynamic compression to a regenerated-cartilage tissue model on the mechanical properties and development of the ECM network.

2. Materials and methods

2.1. Sample preparation and tissue culture

Primary bovine chondrocytes were isolated from metacarpophalangeal joints of steers purchased from a butchery using a sequential enzyme digestion method [10]. Full thickness articular cartilage tissue was harvested from the proximal articular surface of the metatarsal bone and finely diced with a scalpel. The finely diced cartilage tissue was enzymatically digested in a 25 unit/mL protease solution (P8811, Sigma, St Louis, MO) for 3 h and subsequently in a 200 unit/mL collagenase solution (C7657, Sigma) for 18 h at 37 °C. Both enzyme solutions were prepared in sterile tissue culture medium consisting of Dulbecco's modified Eagle's medium (DMEM; D5921, Sigma) supplemented with 20 v/v% Fetal Bovine Serum (FBS; 10437-028, Gibco, Carlsbad, CA), 2 mM L-glutamine (G7513, Wako Pure Chemical Industries, Ltd., Osaka, Japan), 100 unit/mL penicillin, 100 µg/mL streptomycin, 0.25 µg/mL amphotericin B (161-23181, Wako), 20 mM Hepes (H0887, Sigma),

and 0.85 mM L-ascorbic acid (AsA; A5960, Sigma). The supernatant of the resultant solution was centrifuged to separate chondrocytes at 40 × g for 5 min. The resultant cell pellet was washed twice in fresh culture medium, and then cell number and cell viability were determined using Trypan Blue assay. In this study, Sigma Type VII agarose (A6560, Sigma) was used to prepare a chondrocyte-agarose construct. The agarose powder was dissolved in Earle's Balanced Salts Solution (EBSS; Sigma) at twice the required final concentration (1 w/v%) and mixed with an equal volume of the cell suspension to yield the desired agarose concentration with a final cell density of 1×10^7 cells/mL. The molten cell-agarose solution was poured into an acrylic mold and quenched into gel at 4 °C for 30 min to create cylindrical constructs with a diameter of 4 mm and a height of 2.5 mm. The resultant cell-agarose constructs were placed into a 24-well culture plate and cultured with 1 mL culture medium in an incubator at 37 °C and 5% CO₂. The culture medium was exchanged every 2 days. Several culture media with different VC concentrations, from 2.2 to 32 pmol/10⁹ cells, were prepared and used to evaluate the effect of AsA and L-ascorbic acid 2-phosphate magnesium salt (A2P, Wako) concentrations. Culture medium without VC was also used as a control. Each medium is abbreviated as AsA(2.2), A2P(6.4), and A2P(32) hereafter in this paper.

After 1 day of free-swelling culture, the constructs in the experiment group were subjected to uniaxial compression within a purpose-built bioreactor system as shown in Fig. 1. The system permits the application of strain independently in both vertical and horizontal directions to individual constructs using a 24-well plate in a commercially available incubator. The movements were controlled with respective linear variable displacement transducers and linear guide actuators. Strain was applied to the individual constructs through a loading plate, which was attached to the actuator via a jig. Uniaxial cyclic compression up to a maximum amplitude of 15% was applied in a triangular waveform at a frequency of 1 Hz for 6 h and subsequently off-loaded with the plate resting for each construct over the subsequent 18 h. Control constructs were cultured, in contrast, with both upper and lower plates for diffusion through the sides alone.

2.2. Measurement of mechanical property

To examine the influence of VC concentration, cell-agarose constructs were subjected to unconfined compression while immersed in culture medium at room temperature. Individual constructs were tested after culture periods of 22 days. Mechanical tests were

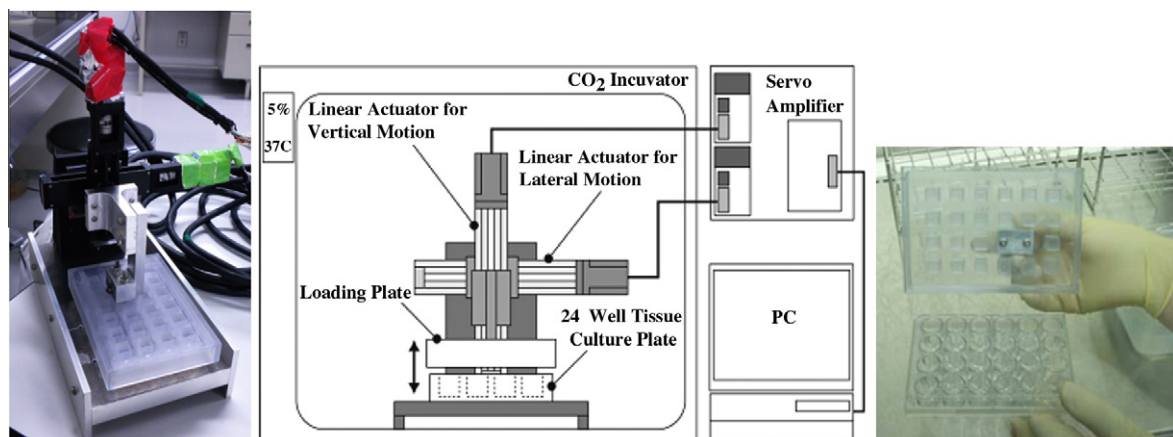


Fig. 1. Photograph (left) and Schematic drawing (center) of mechanical loading system mounted within tissue culture incubator. Loading plate with 22 plungers coupled with a 24-well culture plate (right).

performed with an impermeable stainless steel plunger at a strain rate of 0.5 mm/min up to a strain of 10%, while the load was recorded with a 10 N load sensor. The tangent modulus of the construct was calculated from the slope of a straight-line approximation of the stress–strain curve with a range of 0–15% strain using the least-square method.

2.3. Immunohistology

Separate constructs were used for trichrome immunofluorescence observation to examine the morphological characteristics of the elaborated ECMs, in particular type I collagen, type II collagen and chondroitin sulfate. After the prescribed culture periods, representative constructs were cut into slices with a thickness of approximately 1 mm using a knife. The slices were washed in PBS(–) and subsequently incubated in PBS(–) + 1 w/v% bovine serum albumin (BSA; Wako) for 30 min at 37 °C. These slices were incubated in PBS(–) containing the three monoclonal antibodies (bovine IgG1 isotype anti-type I collagen, Funakoshi, Tokyo, Japan; embryonic chicken IgG2a isotype anti-type II collagen, Funakoshi; mouse IgM isotype anti-chondroitin sulfate, Sigma) for 90 min at 37 °C to primarily label the collagens and the proteoglycan at once. The slices were then washed three times in PBS(–) for 10 min and

incubated in PBS(–) containing the three secondary antibodies corresponding to each of the primary antibodies (Alexafluor 350-conjugated anti-mouse IgG1 antibody, A21120; Alexafluor 488-conjugated anti-mouse IgG2a antibody, A11001; Alexafluor 568-conjugated anti-mouse IgM antibody, A21043, Invitrogen) for 60 min at 37 °C. Labeled ECM molecules were fluorescently visualized within the cultured cell–agarose construct. The fluorescently stained specimens were mounted on the coverslip and observed using a confocal laser scanning microscope (CLSM; Eclipse; Nikon Corp., Tokyo, Japan).

2.4. Cell viability

The cell viability of individual constructs was assessed at different concentration of the VCs using a previously reported protocol [11]. The cultured constructs were cut into slices with a thickness of approximately 1 mm and washed in PBS(–) for 5 min at 37 °C. These slices were incubated in PBS(–) containing two fluorescent dyes (live cells: Calcein AM, Molecular Probes; dead cells: Ethidium homodimer-1, Molecular Probes) for 15 min at 37 °C. The fluorescently stained constructs were then washed by PBS(–) several times and mounted on the coverslip and observed by CLSM.

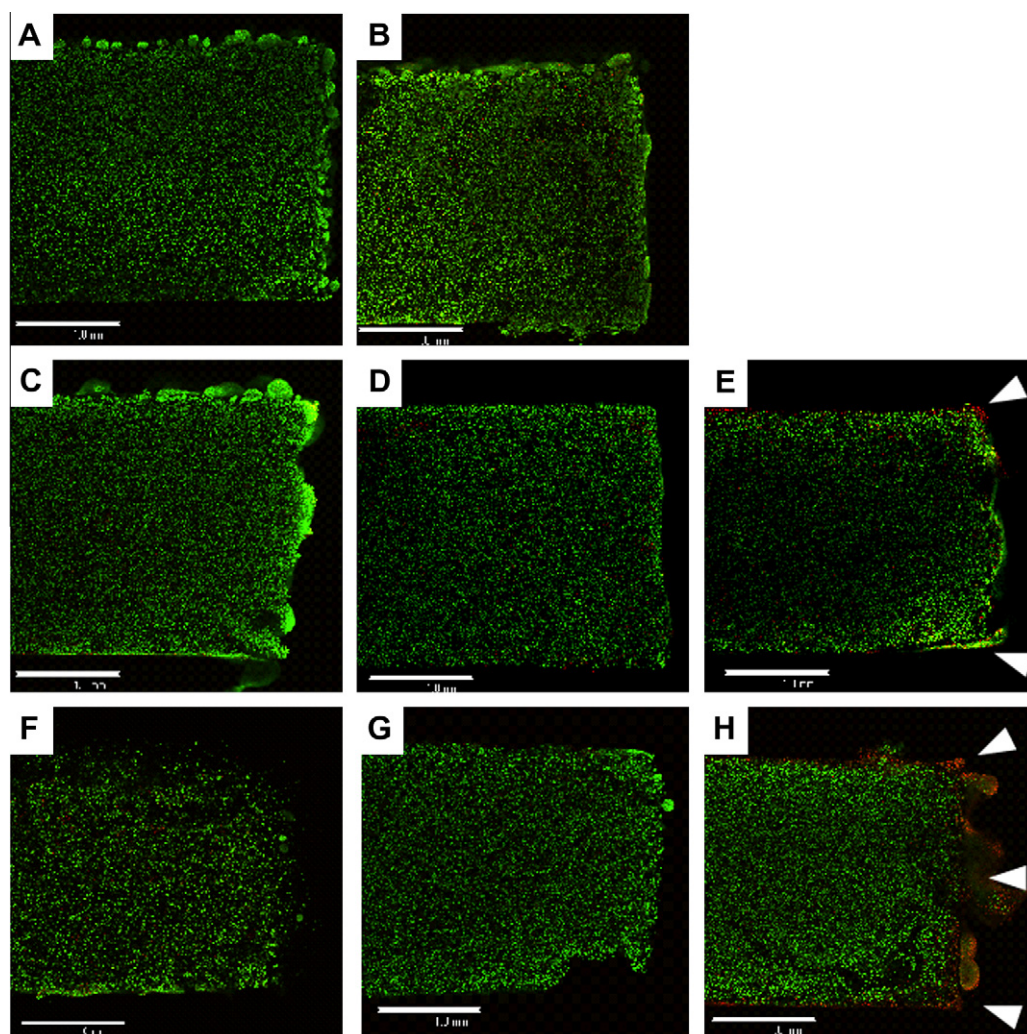


Fig. 2. Low-magnification immunofluorescence images of cell viability under free swelling (A, B, C, F), control (D, G) or compression (E, H) culture conditions. (A): without VC; (B): AsA(2.2); (C–E): A2P(6.4); (F–G): A2P(32). Green represents live cells and red represents dead cells. The culture period was 22 days. Each trichrome-stained sample represents an individual culture. The scale bar represents 1 mm.

2.5. Statistical analysis

Result of the tangent modulus were expressed as the mean \pm sample standard deviation (SD). The significance of the difference between each experimental group was assayed using the two-tailed Welch's *t*-test. The degree of freedom was abbreviated to *df*. If the results of pairwise comparisons between two groups were P -value < 0.01 , $d > 1$ and $(1 - \beta) > 0.8$ simultaneously, we judged the difference as significant in the tangent modulus, where P -value: level of significance; d : Cohen's *d*, an indicator of the effect size [12]; $(1 - \beta)$: the power of the test calculated by R language.

3. Results

3.1. Viability test

In the majority of cases there was a high degree of viability after 22 days of culture, as indicated in Fig. 2. However, some cytotoxicity was evident in the corner of the constructs in two experimental groups, namely A2P(6.4), and most particularly A2P(32), as indicated by the arrow-heads in Fig. 2(E) and (H).

3.2. Measurement of mechanical properties

To evaluate the influence of the mechanical stimulation on the cultured constructs, the tangent moduli of each group after 22 days of culture is shown in Fig. 3. Comparison within the four free-swelling groups only indicated significant differences between the without VC and the AsA(2.2) groups, the latter revealing an increased tangent modulus. By applying cyclic compression to the construct with addition of VC to the culture medium, there was not a clear difference between free-swelling and compression groups for AsA. By contrast, both tangent moduli of the two compression groups were higher than those of several free-swelling samples in each A2P dose group (A2P(6.4) and A2P(32)). Each tangent modulus of the control groups with A2P was ranked against the free-swelling and compression groups. It is not evident that there were differences between each pair of the three compression groups (AsA(2.2), A2P(6.4) and A2P(32)) (AsA(2.2), A2P(6.4):

$df = 16$, $P = 0.47$, $d = 0.32$, $(1 - \beta) = 0.030$), but A2P(32) was lower than both AsA(2.2) and A2P(6.4) because each effect size (d) was large but each power ($(1 - \beta)$) was not (AsA(2.2), A2P(32): $df = 25$, $P = 0.017$, $d = 0.99$, $(1 - \beta) = 0.43$; A2P(6.4), A2P(32): $df = 15$, $P = 0.011$, $d = 1.3$, $(1 - \beta) = 0.54$).

3.3. Histological observation

Fig. 4 shows immunofluorescent images at low and high magnification revealing the ECM distribution (type I and II collagen and chondroitin sulfate) after 22 days. The high-magnification images were taken in the vicinity of the center of the construct. As indicated in Fig. 4(a) and (A), it is clear that with the absence of VC in the culture medium, there was only a limited amount of ECM. By contrast, in the free-swelling group (AsA(2.2)), there was a fairly uniform distribution of ECM across the construct (Fig. 4(b)). As indicated by the arrows in Fig. 4(B), at high magnification, this revealed a collagen network interconnected between chondrocytes. The low-magnification images for each of the A2P dose groups revealed a similar distribution in ECM across the constructs (Fig. 4(c)–(h)), although it was difficult to observe the ECM in the peripheral region of the compressed construct associated with group A2P(32), as indicated by the arrow-heads in Fig. 4(h). In the high magnification images, type II collagen seems to be interconnected between the chondrocytes, as indicated by the arrow in Figs. 4(E), (G) and (H). However, those examinations revealed that the compression groups of A2P(6.4) and A2P(32) were associated with a more abundant collagen network than the corresponding free-swelling groups. In addition, it is clearly evident that the distribution of chondroitin sulfate is restricted to the peripheral regions of the chondrocytes.

4. Discussion

In order to establish a suitable method for regenerating tissue-engineered cartilage, we studied the effects of two types of chemical and physical stimulations on the mechanical properties of the chondrocyte–agarose construct a regenerated-cartilage model. The first stimulation was the addition of high concentration of VC to the culture medium in order to enhance collagen synthesis. The second was the application of a compressive strain to the construct using a purpose-built bioreactor. Our results show that compressive strain had led to an increase in the tangent modulus and that the collagen network had become dense and interconnected among chondrocytes. We have already described the importance of chondrocyte linkage by ECM for the development of bulk elasticity of the construct in a previous study.

It is well known that even though A2P has no physiological activity, it can nevertheless reproduce the effect of VC activity after dephosphorylation by an alkaline phosphatase (ALP) [13]. Dephosphorylated A2P, or AsA, penetrates chondrocytes through a VC transporter, chiefly the sodium-dependent VC transporter 2 (SVCT2), and supports collagen synthesis as a cofactor in the rough endoplasmic reticulum [14]. In free-swelling culture conditions, we only observed a small number of collagen molecules distributed in the construct. We also observed that the tangent moduli of both A2P dose groups were lower than the moduli of AsA(2.2). We must also consider the reaction rates of both dephosphorylation of A2P by ALP and the transport of AsA into the cytosol via SVCT2. The Michaelis constants of bovine chondrocytes ALP and SVCT2 are $1\text{--}10$ and $62 \pm 25 \mu\text{M}$, respectively [15,16]. The affinity of the substrate is slightly higher for ALP than for SVCT2, and as a result, AsA concentration around chondrocytes was lower in the A2P group. The rate of collagen synthesis was also relatively lower in the A2P group than the AsA dose group. Hence, the

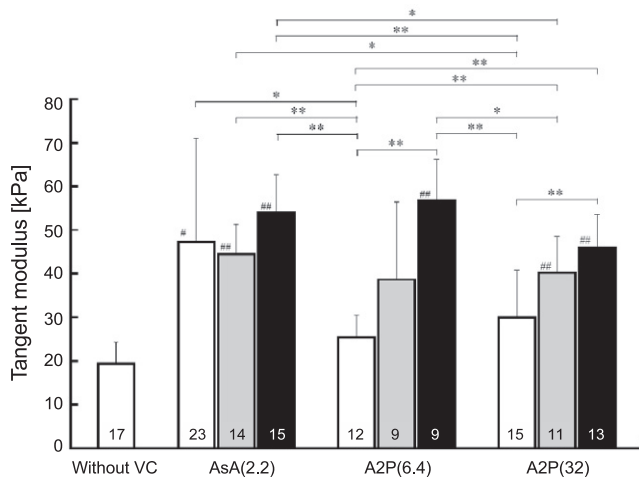


Fig. 3. Comparison of the tangent moduli after 22-day culture period. Cultures were terminated following free-swelling (white column), control (gray), or compression (black) conditions. Each number in the columns is the cultured sample number and each sample represents an individual culture. The error bar represents SD. Sharps (# and ##) and asterisks (* and **) show respective statistical significance between control group and each groups in Welch's *t*-test. #, *: $d > 1$, $P < 0.01$, $(1 - \beta) > 0.8$; ##, **: $d > 2$, $P < 0.01$, $(1 - \beta) > 0.8$.

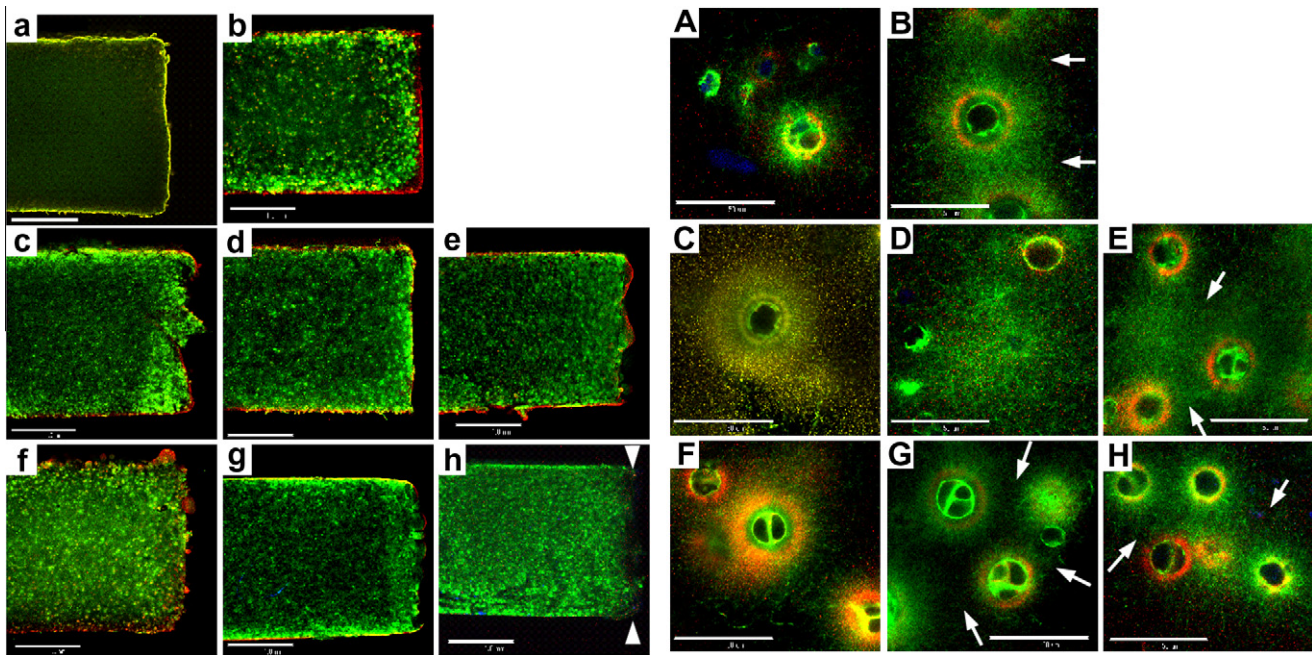


Fig. 4. Low- and high-magnification immunofluorescence images of ECMs under either free-swelling (a, b, c, f, A, B, C, F), control (d, g, D, G), or compression (e, h, E, H) culture conditions. (a, A): without VC; (b, B): AsA(2.2); (c–e, C–E): A2P(6.4); (f–g, F–G): A2P(32). Blue: type I collagen; green: type II collagen; red: chondroitin sulfate. The culture period was 22 days. Scale bars represent 1 mm (a–g) and 50 μ m (A–G), respectively. Arrow-heads indicate that we could not observe ECMs at periphery of the tissue of compression group A2P(32). Arrows indicate interconnections among chondrocytes by type II collagen network.

tangent modulus of the free-swelling groups with A2P was suppressed by restraining cytotoxicity of AsA.

When applying compressive strain to the construct, each tangent modulus of A2P(6.4) and A2P(32) was higher than the moduli for the respective free-swelling groups. This is because the collagen network of these groups expanded, which allowed collagen fibers to interconnect among the chondrocytes. We think this mechanical stimulation enhanced diffusion of A2P and nutrients, homogenizing A2P within the construct. In addition, the stimulation probably excited a mechanosensor, activating cell-signaling pathways, namely the mechanotransduction pathways. It is well-known that mechanical stimulation causes chondrocytes to undergo several biological responses involving cartilage remodeling strategies that are relevant to the implantation of cultured tissue [17], e.g., activation of a mitogen-activated protein kinase (MAPK) pathway, a cell-signaling pathway [18,19], an increase in GAG biosynthesis [20], and regulation of inflammatory species synthesis [21]. These chondrocyte responses followed soon after mechanical stimulation and activity of the MAPK pathway was maintained for 5–60 min [18,19]. In synthesizing ECM with mechanical stimulation, chondrocytes built up the collagen network and adapted to changes in the deformation around the cells. Therefore, the compression group with A2P had a higher tangent modulus than the free-swelling group.

The tangent modulus of the compression group with A2P(32) tended to decrease more than the compression group of A2P(6.4) because ECM had been less developed in the peripheral region of A2P(32) than A2P(6.4). Moreover, viability observations revealed that cytotoxicity occurred at the peripheral regions of A2P(32). Generally, we can think that uniaxial compressive deformation of a cylinder model follows the same physics as two-dimensional tension in a plane perpendicular to the axial direction. When the tangent modulus is uniform in the construct, the force at the outer diameter position will be the largest because a reactive force is directly proportionate to the radius. Therefore, less ECM at the

peripheral region induced a decrease in the tangent modulus of the construct.

As mentioned above, it is not easy to homogeneously develop ECM in free-swelling culture conditions for the purpose of making large-sized regenerated-cartilage tissue. Based on the results of this study, it is necessary to apply mechanical strain to the construct because nutrients should be uniformly supplied. If we also consider the diffusion of nutrients under free-swelling culture conditions, we must note that the diffusion coefficient of water in agarose gel is about $10 \times 10^{-10} \text{ m}^2/\text{s}$ [22,23]. Applying Fick's law, this means that a water molecule diffuses 10 mm per day in the gel. Pluen et al. reported that the diffusion coefficients of proteins in 0.1 M PBS(–) solution, lactalbumin and ovalbumin, were $0.8\text{--}1 \times 10^{-10} \text{ m}^2/\text{s}$ [24], and therefore that nutrient particles diffuse 3 mm per day in the gel. Moreover, if the ECM of the cultured construct is dense, it is difficult for nutrients to diffuse in the tissue and the diffusion coefficient will be decreased. Thus, for clinical implantation of a large-sized regenerated-cartilage tissue in order to treat a cartilage defect, the construct should be prepared by applying mechanical stimulation, which would homogeneously develop the ECM network.

In summary, this study investigated the influence of two types of VC, AsA and A2P, on the development of the ECM network and regulation of the mechanical properties of chondrocyte-agarose constructs. The collagen network of A2P dose groups improved more than the AsA dose group in free-swelling culture conditions and the tangent modulus of the A2P dose groups did not increase. Moreover, it is clear that free-swelling culture conditions suppressed development of ECM of the inner tissue more than the outer tissue. When applying compressive strain to the construct, tangent moduli of the A2P dose groups were higher due to the fact that ECM networks of the inner tissue had been upregulated with interconnections among the chondrocytes. Additionally, we propose that mechanical stimulation enhanced diffusion of nutrients and improved synthesis of the ECM via mechanotransduction

pathways. We conclude that the application of mechanical stimulation to a large-sized engineered-tissue is necessary for the treatment of articular cartilage defects of various sizes and forms.

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