A Photochemical Crosslinking Approach to Enhance Resistance to Mechanical Wear and Biochemical Degradation of Articular Cartilage

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Abstract
Objective. The objective of this study was to evaluate photochemical crosslinking using Al(III) phthalocyanine chloride tetrasulfonic acid (CaSPc) and light with a wavelength of 670 nm as a potential therapy to strengthen articular cartilage and prevent tissue degradation. Design. Changes in viscoelastic properties with indentation were used to identify 2 crosslinking protocols for further testing. Crosslinked cartilage was subjected to an in vitro, accelerated wear test. The ability of the crosslinked tissue to resist biochemical degradation via collagenase was also measured. To better understand how photochemical crosslinking with CASpc varies through the depth of the tissue, the distribution of photo-initiator and penetration of light through the tissue depth was characterized. Finally, the effect of CASpc on chondrocyte viability and of co-treatment with an antioxidant was evaluated. Results. The equilibrium modulus was the most sensitive viscoelastic measure of crosslinking. Crosslinking decreased both mechanical wear and collagenase digestion compared with control cartilage. These beneficial effects were realized despite the fact that crosslinking appeared to be localized to a region near the articular surface. In addition, chondrocyte viability was maintained in crosslinked tissue treated with antioxidants. Conclusion. These results suggest that photochemical crosslinking with CASpc and 670 nm light holds promise as a potential therapy to prevent cartilage degeneration by protecting cartilage from mechanical wear and biochemical degradation. Limitations were also evident, however, as an antioxidant treatment was necessary to maintain chondrocyte viability in crosslinked tissue.

Keywords
photochemical crosslinking, cartilage mechanics, CASpc, wear

Introduction
Osteoarthritis (OA), a leading cause of disability worldwide,¹ is characterized by cartilage degeneration and the eventual loss of cartilage tissue.² In addition, OA cartilage has reduced mechanical properties, including decreased resistance to swelling and diminished elastic modulus in tension, compression, and shear deformations.³⁶ These inferior properties limit the ability of the diseased cartilage tissue to fulfill its mechanical functions of distributing joint loads and providing a low-friction surface that sustains minimal wear.

One method to restore the stability and strength of cartilage is to increase the crosslinks in the collagen network. Collagen has an inherently crosslinked structure, and can obtain additional, non-native crosslinks through exposure to exogenous crosslinking agents. Collagen crosslinking significantly enhances the modulus and strength

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of collagenous tissues, and improves their resistance to enzymatic degradation. Initial studies of cartilage crosslinking used formaldehyde or glutaraldehyde to improve wear properties. However, the crosslinks formed by these chemical agents are toxic to cells. We previously showed that crosslinking with genipin, a natural crosslinking agent with favorable biocompatibility, protects healthy and impact loaded cartilage from mechanical wear and enzymatic degradation. Genipin has also been shown to stabilize decellularized cartilage tissue. Other crosslinking agents, such as lysyl oxidase like-2 and epigallocatechin gallate, also enhance the failure strength and modulus of cartilage. However, despite these potential benefits, it would be challenging to apply any chemical crosslinking agent to damaged or degraded articular cartilage in a patient suffering from OA. Directing the crosslinking agent to a particular tissue or region of tissue would be difficult, and chemical crosslinking agents that are introduced to an articular joint would crosslink not only cartilage but also adjacent tissue within the joint capsule.

A potential alternative crosslinking mechanism is photochemical crosslinking, in which a chemical photosensitizer is excited by light and generates a reaction that covalently bonds collagen fibrils. One advantage of photo-initiated crosslinking is that crosslinking can be localized to a particular region of tissue, such as tissue that has been damaged, by controlling the location of light exposure. Photochemical crosslinking has been demonstrated to crosslink the interface between 2 cartilage surfaces, and may enhance the integration of osteochondral grafts to native cartilage. In particular, Al(III) phthalocyanine chloride tetrasulfonic acid (CASPc), a photosensitizer from the phthalocyanine group that is activated by 670 nm light, was reported to produce functional bonds between 2 cartilage surfaces without significant loss of cell viability. However, whether photochemical crosslinking with CASPc enhances the mechanical properties and resistance to collagenase degradation of cartilage tissue has not been previously investigated.

The objective of this study was to evaluate photochemical crosslinking using CASPc and 670 nm light as a potential therapy to strengthen articular cartilage and prevent tissue degradation. First, changes in viscoelastic properties with indentation were determined for 2 different incubation times in CASPc solution and a wide range of light exposure duration. From these data, 2 different crosslinking protocols were selected for further testing. Crosslinked cartilage was subjected to an in vitro, accelerated wear test to assess the effect of crosslinking on cartilage wear and coefficient of friction (COF). The ability of the crosslinked tissue to resist biochemical degradation via collagenase was also measured. To better understand how photochemical crosslinking with CASPc may vary through the depth of the tissue, the distribution of photo-initiator in the tissue and penetration of light through the depth of the tissue was determined. Finally, the interaction between CASPc and 670 nm light generates singlet oxygen, an oxidant that promotes oxidative damage. Therefore, chondrocyte viability and the effect of an antioxidant treatment with photochemical crosslinking was evaluated.

**Methods**

**Sample Preparation and Photochemical Crosslinking**

Osteochondral specimens were harvested from the condyles of thawed bovine stifles. Cylindrical specimens with 9.5 mm diameter were acquired by coring the condyles such that the articular surface was perpendicular to the coring axis. No more than 3 specimens were obtained per stifle, from locations that provided the flattest articular surfaces. Specimens were stored in gauze dampened with phosphate buffered saline (PBS) at –20°C until the day of testing. Photochemical crosslinking was performed by first incubating thawed specimens in 0.5 mM CASPc (Frontier Scientific, Logan, UT) solution in PBS for 20 or 60 seconds at room temperature. Then, the samples were removed from CASPc solution and exposed to 670 nm light from a fiber-coupled laser (B&W Tech Inc, Newark, DE) for 0, 15, 30, 60, 120, 300, or 600 seconds. During light exposure, clear plastic wrap was placed over the cartilage surface to prevent dehydration. The laser was equipped with a fiber collimator (Edmund Optics, Barrington, NJ) with 0.25 numerical aperture and produced a 4.6 mm diameter beam spot size. To enhance the spatial uniformity of the light intensity, the total fiber length was extended to 180 cm. The power was set such that it was 273.5 ± 0.50 mW after passing through both the fibers and the collimator.

**Viscoelastic Properties from Indentation**

Stress-relaxation indentation tests were conducted on osteochondral specimens with a Mach-1 mechanical tester (Biomometum; Montreal, QC) equipped with a load cell with ±150 g range (AL312AL, Honeywell Industries Inc., Columbus, OH) and a 3.0 mm diameter flat punch indenter with 0.2 mm edge radius. During indentations, specimens were submerged in a hydrating solution consisting of PBS with protease inhibitors (1 mM ethylenediaminetetraacetic acid, 5 mM benzamidine, and 10 mM n-ethylmaleimide). Stress-relaxation tests started with a load threshold of 0.0875 gm force to find the contact at the cartilage surface and then consisted of a 20-second linear ramp to reach 50 µm peak displacement, followed by a 20-second hold at peak displacement. Five locations on each specimen were indented, with the first location near the center of the specimen and the other 4 locations 1 mm from the center. The
stress-relaxation indentation was performed in quadruplicate at each location, then specimens were crosslinked with CASPc, as described above, and the indentation tests were repeated at the same 5 locations. The load-time data during stress relaxation was utilized to find 3-dimensional best-fit parameters $E_1$, $E_2$, and $\eta$ of a standard linear solid (SLS) model, which has a parallel spring and dashpot in series with a free spring, using MATLAB (Mathworks, Natick, MA) optimization toolbox.\textsuperscript{11,20} Instantaneous equilibrium ($E_{eq}$) and equilibrium ($E_{eq}$) modulus and time constant ($\tau$) due to crosslinking were calculated from the best-fit parameters. Values for the quadruplicate indentations at each location before and after crosslinking were averaged. The effect of crosslinking was evaluated at each location from the ratio of SLS parameters after crosslinking normalized by the value of the same parameter before crosslinking ($n = 10-15$ locations). From these results, 2 crosslinking protocols were selected for further study: (1) 20 seconds in CASPc solution and 600 seconds light exposure, denoted 20C600L, and (2) 60 seconds in CASPc solution and 30 seconds light exposure, denoted 60C30L.

**Wear and Friction**

Thawed osteochondral cores were crosslinked using one of the two selected protocols ($n = 7$ for each protocol). Untreated specimens were exposed to neither CASPc nor light as controls ($n = 10$). We previously reported differences in wear performed perpendicular or parallel to the preferential fiber direction.\textsuperscript{21} Therefore, the preferential direction of the collagen fibers was determined with a split line analysis. The cartilage surface was pierced with a pin at 10 to 12 locations on the periphery. India ink was applied to the surface of the specimen and cleaned immediately with gauze. The split lines revealed the preferential direction of collagen fibers at the articular surface.\textsuperscript{22}

An accelerated wear test was performed with a reciprocating motion against a T316 stainless steel plate (#8 mirror finish) using a UMT Tribolab (Bruker, San Jose, CA) equipped with a 2-axis load cell. The specimen was manually oriented such that the split lines were aligned with the direction of wear. The specimen was loaded to a constant force of 160 N. This load was maintained for 14,000 reciprocating cycles of 18 mm length in each direction at a speed of 4 mm/second with 1 second pause at each end.\textsuperscript{21} In addition, untreated specimens that were exposed to the reciprocating motion of the wear test without loading were included to better understand the effect of the mechanical load ($n = 3$). The 43.5 hours wear test was conducted at room temperature in PBS containing protease inhibitors.

Wear was measured by biochemically quantifying the amount of sulfated glycosaminoglycans (sGAGs) and hydroxyproline (HYP) that were released to the solution bath during the wear test. The solution bath was collected at the end of each experiment, lyophilized, and rehydrated with water. The solution was filtered using ultra centrifugal filter (Amicon\textsuperscript{8}, 3000 MWCO, Millipore Corporation, Billerica, MA) at 4,000 g acceleration for 90 minutes. Preliminary studies found that removing salts and protease inhibitors from the solution via filtering provided more repeatable biochemical measurements, and that matrix components were not detected in the filtrate. The filtered wear particles were digested with papain, and sGAG content was measured using the dimethyl-methylene blue (DMMB) assay.\textsuperscript{21,23,24} Separate aliquots were assessed for HYP content with a chloramine-T assay.\textsuperscript{23-26} The amount of collagen in the solution was calculated based on a HYP: collagen ratio of 1:7.69.\textsuperscript{27} The quantities of matrix components that were released from the specimens that were not loaded were subtracted from the respective values from the wear test to determine the matrix components released due to mechanical wear only.

Friction and normal loads were recorded by UMT Tribolab load sensor at a rate of 100 Hz during the wear test and initial and equilibrium COF were extracted. For calculating COF, MATLAB (Mathworks, Natick, MA) code was developed to discard data from the acceleration and deceleration of each cycle so that only data at 4 mm/second velocity was considered. An average value of COF was calculated for each reciprocating cycle. The averaged value for the first reciprocating cycle was considered the initial COF and the average of last 8,000 cycles was considered the equilibrium COF.\textsuperscript{21}

**Collagenase Digestion**

The effect of photochemical crosslinking on the resistance of cartilage to degradation by collagenase was investigated. Osteochondral specimens were treated with one of the two selected crosslinking protocols or were left untreated ($n = 5$ for each protocol). Using a sledge microtome (HM 450 Richard Allan, Kalamazoo, MI) equipped with a freezing stage (Physitemp, Clifton, NJ), three 50 $\mu$m sections were taken starting from the articular surface of each specimen. Individual sections were incubated for 45 minutes at 37 °C in 0.5 ml of a 2 mg/ml solution of type I collagenase from Clostridium histolyticum in 50 mM Trizma buffer at pH 7.4 containing 10 mM CaCl$_2$ (all from Sigma-Aldrich, St. Louis, MO).\textsuperscript{11,12} To quantify the digested collagen, the digest solution for all sections were collected and assayed for HYP and the amount of digested collagen was estimated, as above.

**CASPc and Light Penetration**

To observe CASPc penetration into the cartilage and the effect of light exposure, osteochondral specimens were incubated in CASPc for 20 seconds, with and without 600
seconds light exposure, or for 60 seconds incubation, with and without 30 seconds of light exposure. Then, specimens were cut in half with a scalpel, and the cross section was imaged with a stereomicroscope (Stemi 508, Carl Zeiss, Jena, Germany).

To study light penetration through the cartilage depth, including through cartilage that was treated with CASPc, specimens were incubated in CASPc solution for 20 or 60 seconds or were left untreated ($n = 3$ for each group) and then four 50-µm thick sections were cut starting from articular surface as described previously for collagenase digestion (Section “Collagenase Digestion”). Transmission of 670 nm light from a laser through each section was measured. Each hydrated section was placed in the center of a plastic shim in the shape of a ring, and both the section and shim were sandwiched between a glass slide and cover slip. The plastic ring shim had the same thickness as cartilage sections (50 µm) and acted as a spacer to prevent the tissue from being compressed. All 4 sections of both treated and control samples were exposed to 670 nm light for 600 seconds while the power of light transmitted through the tissue was measured and recorded using a light sensor (PowerMax-USB, Coherent Inc, Santa Clara, CA) at 10 Hz. The power of light transmitted through the glass slide and cover slip without cartilage was also determined.

Chondrocyte Viability and Effect of Antioxidants

Fresh metacarpophalangeal joints were obtained from a local abattoir (Kenny’s Fine Meats, Mooresville, IN). Cartilage tissue was extracted using a 6 mm biopsy punch and sterile scalpel using aseptic techniques. The cartilage explants were cultured in Dulbecco’s modified Eagle medium (DMEM) containing high glucose and L-glutamine (Corning Inc., Corning, NY) supplemented with 50 µg/ml ascorbic acid (Sigma-Aldrich) and Insulin-Transferrin-Selenium (Gibco, Grand Island, NY). For the antioxidant treatment, 20 mM N-acetyl-L-cysteine (NAC; Sigma-Aldrich) and 10 mM sodium pyruvate (Sigma-Aldrich) were added to the culture medium and CASPc solution. After 24 hours in culture, the explants were aseptically crosslinked with the 60C30L crosslinking protocol with or without antioxidants added to the CASPc solution or were incubated in CASPc solution alone for 60 seconds but were not exposed to 670 nm light as a control. After crosslinking, explants were rinsed and were cultured either with or without antioxidants as per their initial treatment for an additional 24 hours.

To determine the effect of crosslinking and antioxidants on chondrocyte viability, explants were cut in half, and incubated with 2.5 µM calcein AM to stain the live cells green and 5 µM ethidium homodimer to stain the dead cells red (LIVE/DEAD™ Viability/Cytotoxicity Kit, Invitrogen, Carlsbad, CA) for 30 minutes. After rinsing in PBS, specimens were imaged using confocal fluorescent microscopy (Olympus FV-1000 MPE). To evaluate apoptotic signaling, additional explant halves were stained with CellEvent caspase 3/7 reporter (Invitrogen) and imaged.

In addition, whether antioxidants inhibited the photochemical crosslinking reaction was determined by evaluating viscoelastic properties of cartilage that had been crosslinked with or without antioxidants. The indentation test was performed on specimens from thawed condyles before and after the 60C30L crosslinking protocol with and without the addition of antioxidants in the CASPc and PBS solutions ($n = 10-15$).

Statistics

Differences between crosslinking treatments were determined using one-way and two-way analyses of variance (ANOVA) with Dunnett’s post hoc with significance set at $p < 0.05$ (OriginPro 2020, OriginLab Corporation, Northampton, MA). Data are presented as mean ± standard deviation.

Results

Viscoelastic Properties of Cartilage

The instantaneous stiffness, equilibrium stiffness, and relaxation time constant in untreated tissue were 3.13 ± 1.28 MPa, 0.37 ± 0.15 MPa, and 0.92 ± 0.09 seconds, respectively. The instantaneous and equilibrium modulus of cartilage was increased by photochemical crosslinking, and was dependent both on the incubation and light exposure times. In both 20 and 60 second CASPc incubation times, the average instantaneous modulus increased by at least 4.8% at higher light exposure times (120-600 seconds; Fig. 1A and D). With 20 seconds incubation in CASPc solution, the equilibrium modulus also showed increases of 14.8% to 22.3% with light exposure of more than 120 seconds (Fig. 1B). Equilibrium modulus with 60 seconds incubation in CASPc solution alone for 60 seconds but were not exposed to 670 nm light as a control. After crosslinking, explants were rinsed and were cultured either with or without antioxidants as per their initial treatment for an additional 24 hours.

Wear and COF

Cartilage wear was quantified by the amount of cartilage matrix components released to the fluid bath during the wear test due to the mechanical load. The amount of collagen and sGAGs released into the solution bath for specimens that were not loaded was 69.3 ± 32.4 µg and 1.92 ±
0.47 mg, respectively. These values were subtracted from the respective values from the wear test to determine the amount of matrix components released because of mechanical wear only.

The average amount of collagen released into the fluid bath due to mechanical wear for the untreated controls was 0.54 ± 0.23 mg. For the 2 selected crosslinking protocols, 20C600L and 60C30L, 0.52 ± 0.28 mg and 0.55 ± 0.22 mg were released, respectively (Fig. 2A). No significant difference in collagen released during the wear test was detected between controls and crosslinked specimens. For the untreated control group, the amount of sGAGs released due to wear was 5.88 ± 4.17 mg. The sGAG wear decreased to 2.04 ± 1.12 mg and 1.69 ± 1.63 mg in the 20C600L and 60C30L crosslinked groups, respectively (Fig. 2B).

For both treated and untreated samples, the COF started from a low value and increased with time, similar to previous reports. The initial COF of control specimens was 1.59 ± 0.63 × 10⁻³ and for the 20C600L and 60C30L crosslinking protocols, it was 1.42 ± 0.65 × 10⁻³ and 1.27 ± 0.53 × 10⁻³, respectively. No significant difference was detected between crosslinked and control specimens for initial COF. Equilibrium COF for controls was 0.21 ± 0.03 and for crosslinking protocols, 20C600L and 60C30L, its values were 0.26 ± 0.03 and 0.24 ± 0.05, respectively (Fig. 3). The equilibrium COF of the 20C600L specimens tended to be larger than controls (p = 0.058).
Collagenase Digestion

The effect of photochemical crosslinking on cartilage’s resistance to collagenase digestion was assessed. The amount of collagen digested from the most superficial section treated with 60 seconds CASPc incubation and 30 seconds light exposure was slightly smaller than that of controls, while there was no difference in the amount of collagen released in other sections from the crosslinked groups compared with control (Fig. 4).

Light Attenuation and CASPc Penetration

The power of transmitted light through 50 µm sections of cartilage was measured for 600 seconds to study the effect of CASPc on the light transmission through cartilage. The power of transmitted light through the microscope slides without cartilage was 257.1 ± 0.3 mW. This power decreased to 199.2 ± 0.1 mW when light was transmitted through the most superficial section of the controls. The power of transmitted light was approximately constant.
over time in all sections of the untreated controls (Fig. 5A). For the most superficial section treated with 20 seconds CASPc incubation, the power of transmitted light started from a lower value of 153.6 ± 17.5 mW and increased to 186.6 ± 9.4 mW after 600 seconds (red curve in Fig. 5B). The same behavior was observed for 60 seconds CASPc incubation time, except the transmitted light’s power started from a lower value compared with the 20 seconds CASPc incubation time (116.7 ± 35.8) before reaching its maximum value at 600 seconds (178.7 ± 6.7 mW, red curve in Fig. 5C). For sections taken below the articular surface, the power of transmitted light was approximately constant over time and similar to the control values (Fig. 5).

As CASPc is highly pigmented, the relative amount of CASPc in the tissue is distinguishable by the intensity of color. After 20 and 60 seconds incubation in CASPc, all specimens become aqua-blue in color, with the longer incubation time in CASPc leading to a darker color (Fig. 6, first row). In addition, the most intense color in both incubation times was at the surface of the specimen, with very little indication of the photochemical initiator deeper in the tissue. Exposure of the tissue to 670 nm light reduced the color intensity at the location of light exposure. This was most evident with 20 seconds of CASPc incubation (Fig. 6, second row).

**Chondrocyte Viability and Effect of Antioxidants**

Chondrocyte viability was maintained in specimens incubated in CASPc solution alone for 60 seconds, in the absence of 670 nm light exposure. In contrast, a band of dead chondrocytes (stained red) was observed at the articular surface in specimens crosslinked with the 60C30L protocol that were not treated with antioxidants (Fig. 7). When antioxidants were added to the culture medium and CASPc solution, chondrocyte viability in crosslinked specimens (Fig. 7) was similar to specimens that were not crosslinked. Consistent with these results, apoptotic signaling via the caspase 3/7 pathway was elevated in crosslinked specimens compared with those that were only exposed to CASPc and was rescued with the addition of antioxidants (Fig. 7).

The effect of the antioxidant treatment on photochemical crosslinking was evaluated. Viscoelastic properties were determined from stress-relaxation indentation before and after crosslinking with the 60C30L protocol with and without the addition of antioxidants. Moduli did not change with the antioxidant treatment (Fig. 8). However, the relaxation time constant increased by an average of 10% with the addition of antioxidants.
Discussion

This study investigated 2 photochemical crosslinking protocols using CASPc and 670 nm light to strengthen the collagen network of cartilage. The first crosslinking protocol, denoted 20C600L, consisted of 20 seconds incubation in CASPc solution and 600 minutes light exposure. This protocol was chosen because it was previously reported to adhere two cartilage surfaces to one another.\(^\text{19}\) A longer

CASPc incubation of 60 seconds was also tested, and resulted in increased CASPc in the tissue, as evidenced by increased light attenuation through sections taken from the articular surface (Fig. 5) and a more intense color of the photo-initiator (Fig. 6). The higher level of CASPc in the tissue resulted in a significant increase in indentation modulus with as little as 30 seconds light exposure (Fig. 1). Therefore, a second protocol consisting of 60 seconds incubation in CASPc solution and 30 seconds light duration (denoted 60C30L) was also selected for further study. The 30 seconds duration of light exposure coupled with 60 seconds of CASPc treatment produced protective effects on articular cartilage that were similar to those of the much longer 600 seconds duration of light exposure following 20 seconds of CASPc treatment. This shorter duration of light...
exposure may be more feasible to implement in a clinical setting than a longer one. In addition, minimizing the duration of light exposure may limit exposure to oxidants and any other potential adverse effects. Results suggest that photochemical crosslinking holds promise as a potential therapy to prevent cartilage degeneration, as crosslinked tissue released less sGAGs during the in vitro wear test (Fig. 2). Resistance to collagenase digestion also improved with crosslinking, although the effect was modest (Fig. 4). Potential limitations were also evident, however, as an antioxidant treatment was necessary to maintain chondrocyte viability and suppress apoptotic signaling in crosslinked tissue (Fig. 7).

Results suggest that photochemical crosslinking with CASPc is limited to a region near the articular surface that is not more than 50 µm deep. The pigmented color of the photochemical initiator remained near the articular surface and did not appear to penetrate deep into the tissue (Fig. 6). Furthermore, 670 nm light interacts with CASPc to induce the formation of singlet oxygen,19 which is necessary for the crosslinking reaction. Singlet oxygen is very reactive and unstable, generating hydrogen peroxide and free oxygen radicals. These molecules are implicated in damaging cellular mechanisms including apoptosis, inflammation, aging, and the injury response.37,39 Sodium pyruvate, which scavenges hydrogen...
peroxide,\textsuperscript{40,41} and NAC, which scavenges free oxygen radicals,\textsuperscript{42} were added to the CASPc solution and media to protect against oxidative stress. The results of the current study indicate that an antioxidant treatment maintains cell viability and reduces apoptosis, although the minimum concentrations and durations of the antioxidant treatment have yet to be determined. Importantly, the antioxidant treatment did not interfere with the photochemical crosslinking reaction, as increases in modulus via indentation with crosslinking were maintained with the antioxidant treatment (Fig. 8).

In conclusion, results of this study indicate that CASPc photo-initiator and 670 nm light strengthen articular cartilage by crosslinking collagen in the superficial zone of the tissue. Parameters of the crosslinking protocol, such as the duration of light exposure and the amount of the chemical photo-initiator in the tissue, were interrelated. These data suggest that specific combinations of photo-initiator concentration, incubation time, light intensity, and light exposure duration may be optimized in future studies to maximize desirable outcomes. The crosslinking treatment at the articular surface was sufficient to reduce the loss of sGAGs from the tissue during an accelerated in vitro wear test. Although crosslinking increased the cartilage’s resistance to biochemical degradation via collagenase, this effect was minimal and was only observed at the articular surface. In spite of these potential benefits, challenges remain before photochemical crosslinking with CASPc can be tested in vivo as a treatment to reduce cartilage degradation. In particular, antioxidants were necessary to preserve chondrocyte viability in crosslinked cartilage. More work is necessary to determine effective antioxidant treatments that maintain chondrocyte health during photochemical crosslinking procedures. In addition, the effect of crosslinking on fluid flow and the exchange of nutrients and waste products in the tissue has not been studied. Furthermore, whether the altered mechanical environment of the tissue influences chondrocyte behavior requires further understanding.

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Ethical Approval
The authors attest that all animal tissue used in this study was collected from a local abattoir. No human tissue was used in this study.

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