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Key Words
Synovial joint · Cartilage · Plowing · Biomechanics · Biochemistry · Chondrocytes · Gene expression · Extracellular matrix

Abstract
Mechanical stress is listed as a main risk factor for cartilage degradation. The aim of this study was to investigate the biological response of cartilage to dynamic loading such as plowing. Cartilage strips harvested from bovine nasal septum were submitted to plowing using a cylindrical indenter, applying a constant normal force in the vertical axis and moving at constant speed in the horizontal axis. After plowing, cell viability, gene expression and glycosaminoglycan (GAG) release were measured with conventional assays. The cell-viability assay and qRT-PCR showed that plowing induces cell death and matrix metalloproteinase 3 (MMP-3) upregulation. The addition of actinomycin D, before or after plowing, confirmed that plowing was responsible for the observed MMP-3 upregulation. Even if the transcriptions of the tissue inhibitor of metalloproteinase (TIMP-1), aggrecan (Agg), collagen type I (Coll1), collagen type II (Coll2) and fibronectin (Fn) were not significantly affected by plowing, actinomycin D treatment revealed that plowing induces a strong increase in TIMP-1 and Coll1 messenger RNA content and influences the gene regulation of Agg, Coll2 and Fn. Furthermore, plowed cartilage explants exhibited enhanced GAG release. Application of hydroxamate MMP inhibitor after loading showed that plowing induces GAG release via the activation of catabolic enzymes. Plowing causes cell death of the chondrocytes closer to the surface as well as matrix damage, observed as GAG loss. Moreover, in healthy chondrocytes, plowing promotes the production and activation of catabolic enzymes like MMP-3.

Abbreviations used in this paper
\begin{tabular}{|ll|}
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18S-rRNA & 18S ribosomal RNA \\
Agg & aggrecan \\
Coll1 & collagen type I \\
Coll2 & collagen type II \\
DMEM & Dulbecco’s modified Eagle’s medium \\
Fn & fibronectin \\
GAG & glycosaminoglycan \\
GAPDH & glyceraldehyde 3-phosphate dehydrogenase \\
GM6001 & hydroxamate matrix metalloproteinase inhibitor \\
MMP-3 & matrix metalloproteinase 3 \\
mRNA & messenger RNA \\
TIMP-1 & tissue inhibitor of metalloproteinase \\
TMJ & temporomandibular joint \\
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Pristine Cartilage

Introduction

Over time, mechanical stress is one of the causes of cartilage degradation. In order to understand the pathomechanics of cartilage breakdown that occurs in the synovial joints, the response of cartilage to mechanical loading has been extensively studied by means of several laboratory models [Kurz et al., 2005]. Interestingly, it has been reported that moderate loading helps cartilage homeostasis [Lane et al., 2000; Griffin and Guilak, 2005; Dossunbekova et al., 2007; Torzilli et al., 2010] but that injurious overload contributes to or causes cartilage degradation [Patwari et al., 2001; Chen et al., 2003; Patwari et al., 2003; DiMicco et al., 2004; Lin et al., 2004; Lee et al., 2005; Sauerland and Steinmeyer, 2007; Verteramo and Seedhom, 2007; Ding et al., 2010]. Chen et al. [2003] showed that continuous or intermittent uniaxial loads of varying durations and magnitudes applied to bovine articular cartilage explants caused chondrocyte death and collagen damage. DiMicco et al. [2004] reported that uniaxial unconfined compression of bovine articular cartilage explants caused proteoglycan degradation and the subsequent loss of glycosaminoglycans (GAGs). Moreover, uniaxial cyclic loading of cartilage explants leads to cell death, collagen damage and GAG loss. It has also been shown that long-term mechanical stress causes an increase in the activity of stromelysin 1/matrix metalloproteinase 3 (MMP-3) [Lin et al., 2004]. Other groups [Lee et al., 2005; Fitzgerald et al., 2006; Sauerland and Steinmeyer, 2007; Verteramo and Seedhom, 2007; Ding et al., 2010] have reported that static compression or shear stress regulates clusters of functionally related gene patterns. Although these in vitro models reproduce the mechanical stresses that cartilage experiences in vivo and provide insights into the biological response of such a complex tissue to mechanical injuries, their uniaxial design presents some limitations.

In a previous study, we studied the temporomandibular joint (TMJ) disk deformations that occur during mandibular function by means of novel, 3-dimensional modeling software that processes data acquired by magnetic resonance imaging and jaw tracking (a system that allows observation of mandibular movement dynamically) [Pallal et al., 2003; Gallo, 2005; Gallo et al., 2006]. These TMJ disk measurements were used to develop a rolling/plowing explant test system that is able to mimic the in vivo plowing effect that results from the combination of compression and sliding of the mandibular condyle on the TMJ cartilage disk [Colombo et al., 2011].

We report here on the biological response of bovine nasal septum cartilage to plowing, by studying chondrocyte viability, variations in gene expression and GAG release.

Materials and Methods

Harvesting of Cartilage Explants

The nasal septa of 12-month-old calves were provided by a local abattoir within 4 h of slaughter. Under sterile conditions, control cartilage explants (20 × 30 × 2 mm) and cartilage strips (60 × 17 × 2 mm) were harvested by using a ‘dual-parallel-blade cartilage cutter’. Control samples and cartilage strips were washed in Dulbecco’s PBS (Invitrogen, Carlsbad, Calif., USA) and equilibrated overnight in Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen) supplemented with 10% fetal bovine serum (Sigma, St. Louis, Mo., USA), 10 mM HEPES (Sigma), nonessential amino acids (0.1 mM), penicillin (100 units/ml), streptomycin (100 μg/ml) and amphotericin B (0.25 μg/ml).

Plowing of Cartilage Strips

Before plowing, the cartilage strips were glued at their ends to Plexiglas supports with cyanoacrylate glue. The plowing was performed unidirectionally in DMEM for 2 h at 37°C in a sterile hood, using a stainless-steel, nonrotating, cylindrical indenter (25 mm in diameter), moving in the horizontal axis with a constant speed of 10 mm/s and simultaneously compressing the cartilage in the vertical axis by applying normal forces of 25, 50 or 100 N. The indenter stroke length was 40 mm and the total number of plowing cycles was 500. During plowing, the control cartilage was placed as a free-swelling explant in the medium in which the cartilage strip was located.

After plowing, using a blade, the cartilage subexplants (15 × 15 × 2 mm) were collected from both the plowed strip and the free-swelling control, and were subjected to analysis.

Chondrocyte Viability Assay

Following plowing (at 25, 50 or 100 N of applied normal force), the cartilage was sliced into 500-μm sections, rinsed in Dulbecco’s PBS and incubated for 20 min in DMEM containing 1 μg/ml calcine acetoxyethyl ester (live-cell staining; Sigma) and 1 μM ethidium homodimer (dead-cell staining; Sigma). The treated slices were thereafter transferred into fresh DMEM. Fluorescence-microscopy images were acquired at 515 and 635 nm by means of an inverted fluorescence microscope (Zeiss, Axiosvert 200) equipped with a digital camera.

RNA Extraction and qRT-PCR

After plowing at 25, 50 and 100 N of applied normal force, the cartilage subexplants collected from the plowed strips and from the controls were equilibrated for 2, 4 or 24 h in DMEM at 37°C. Upon equilibration, samples were snap-frozen in liquid nitrogen and stored at −80°C. RNA extraction was performed according to Davidson et al. [2006] with some modifications. Finely sliced cartilage subexplants (approx. 50 mg) were placed in Eppendorf tubes and

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homogenized twice for 1 min in 800 μl of TRIzol reagent (Invitrogen). After 5 min of equilibration at room temperature, 200 μl of chloroform was added and the tubes were vigorously shaken, mixed and incubated for 2 min at room temperature. Following centrifugation at 9.5 g for 30 min at 4°C, the obtained aqueous phases were recovered, extracted with 200 μl of chloroform and treated as previously described. The recovered supernatants were transferred into 2-ml tubes, gently mixed with 500 μl of isopropanol, incubated for 10 min at room temperature and subsequently centrifuged at 9.5 g for 40 min at 4°C. The supernatants were discarded and the pellets were resuspended in 900 μl of lysis buffer (RNeasy mini kit; Qiagen GmbH, Hilden, Germany) supplemented with 90 μl of β-mercaptoethanol (Sigma-Aldrich). After adding 900 μl of ethanol (75%), the RNA was purified using an RNeasy mini kit, while genomic DNA was digested with a DNase kit (Qiagen) according to the manufacturer’s instructions. Reverse transcriptions of RNA were performed with random hexamer primers using a 1st-strand cDNA synthesis kit for RT-PCR with avian myeloblastosis virus (Roche Diagnostics AG, Rotkreuz, Switzerland) according to the manufacturer’s instructions. qRT-PCR of the obtained complementary DNA was performed in 96-well plates by means of an iCycler real-time detection system® (iQ5Bio-Rad Laboratories, Hercules, Calif., USA) and the reactions were carried out using a Quantifast® SYBR® Green PCR kit (Qiagen). The primers used were the same as those reported in Fitzgerald et al. [2006]. They were designed for the amplification of selected target genes: the extracellular matrix proteins aggrecan (Agg), collagen type I (Coll1), collagen type II (Coll2), fibronectin (Fn), the catabolic enzyme MMP-3 and its inhibitor [the tissue inhibitor of metalloproteinase (TIMP-1)] and the housekeeping genes glyceraldehyde 3-phosphate dehydrogenase (GAPDH) and 18S ribosomal RNA (18S-rRNA). Housekeeping genes as controls from treated samples and a gene of interest from untreated control samples were run on every plate for normalization purposes. The gene regulation was calculated as a multiple, by using the comparative threshold cycle (ΔΔCt) method. Thus, a gene was considered to be upregulated when the multiple of expression was >2 and downregulated for values <0.5.

**GAG Release**

In order to measure the GAG release following plowing, cartilage subexplants were collected from strips plowed at 100 N of applied normal force and from free-swelling controls. The samples were incubated in DMEM at 37°C in the course of 4 different periods of equilibration (i.e. over 1, 2, 3 or 4 days) after plowing. At each time point, cartilage subexplants and 1 ml of the corresponding culture medium were collected and stored at –20 °C until a period of equilibration (i.e. over 1, 2, 3 or 4 days) after plowing. At each time point, cartilage subexplants and 1 ml of the correspond-

**Inhibition Experiments**

To confirm that the change of chondrocyte gene expression was induced by plowing, 2 different experiments were performed.

(1) Cartilage strips and controls were incubated in DMEM supplemented with 30 μM transcription inhibitor actinomycin D (Sigma) for 2 h before plowing. The strips were then plowed for 2 h with 100 N of applied normal force. In this experiment, the effect of plowing on gene expression was blocked so that no difference with regard to the controls was expected if gene expression changes were due to mechanical loading. (2) Conversely, cartilage strips were plowed for 2 h with 100 N of applied normal force and then incubated for 2 h with the transcription inhibitor. The control cartilage was subjected to the same treatment in the inhibitor. In this case, we would detect only those genes expressed during plowing. After both experiments, gene expression of MMP-3, TIMP-1, Coll1, Agg, Fn and Coll2 was determined by qRT-PCR.

To confirm that plowing causes GAG release by activating catabolic enzymes, experiments of MMP inhibition were performed by using hydroxamate MMP inhibitor (GM6001; Millipore, Bedford, Mass., USA). Briefly, cartilage strips and free-swelling controls were incubated for 4 h before plowing in serum-free DMEM supplemented with 1% SITE (sodium selenite, insulin, transferrin and ethanolamine; Sigma) and containing 10 μM GM6001. After 2 h of plowing at 100 N of applied normal force performed in serum-free medium, cartilage subexplants (3 for each strip) were collected from both control cartilage and the plowed strips and again incubated in DMEM containing GM6001 and SITE for 1, 2, 3 or 4 days.

**Statistical Analysis**

All experiments were carried out in triplicate, each time from a different nasal septum. Results are expressed as the mean ± SE. Statistical differences were analyzed using two-way ANOVA and the Student’s t-test.

**Results**

**Cell Viability**

After plowing the cartilage strips, chondrocyte viability was studied by means of calcein acetoxymethyl ester and ethidium homodimer staining (fig. 1). A layer of dead cells was detected at the surface of the explants. The layers varied in thickness with the magnitude of the applied normal force: they corresponded to 4.6 (p < 0.01), 7.3 (p < 0.01) and 8.7% (p < 0.001) of the total sample thickness for strips plowed at 25, 50 and 100 N, respectively.

**Effect of Plowing on Cartilage Gene Transcription after Equilibration Conducted after Plowing**

Upon plowing completion at 25, 50 or 100 N, cartilage explants from plowed strips and relative controls were equilibrated for 2, 4 or 24 h, and at each time point, the effect of plowing on expression of the selected genes (MMP-3, Fn, Coll1, Coll2, Agg and TIMP-1) was measured using qRT-PCR (fig. 2).
After 2 h of equilibration, cartilage explants subjected to the 25-newton plowing revealed only a slight change in the MMP-3 messenger RNA (mRNA) synthesis (1.8-fold upregulation) and this value increased (2.3-fold) after 4 h but then dropped to 0.9-fold after 24 h.

Furthermore, the cartilage strips subjected to the 50-newton plowing had a 4.7-fold upregulation of MMP-3 after 2 h of equilibration, and this value decreased over time (4.1-fold after 4 h and 2.4-fold after 24 h).

The upregulation of MMP-3 for the cartilage strip subjected to the 100-newton plowing was stronger: 6.3-fold after 2 h, 5-fold after 4 h and 3.2-fold after 24 h of equilibration conducted after plowing.

In general, the decrease in MMP-3 expression was statistically significant, comparing the expression after 2 and 24 h for the 25-newton and 100-newton applied normal forces (for 25 N: p = 0.0001, for 50 N: p = 0.0700 and for 100 N: p = 0.0075).

The plowing at the chosen applied normal forces did not influence the net gene expression of Agg, Fn, TIMP-1, Coll1 and Coll2, the levels of which remained at around the preplowing values (fig. 2).

Transcription Inhibition Experiments
Upon plowing completion and without inhibitor treatment, MMP-3 was upregulated 8.6-fold (fig. 3a), but when the cartilage strips were treated with actinomycin D for 2 h before plowing, no MMP-3 upregulation was observed upon plowing completion (p = 0.0001). In contrast, the treatment with the transcription inhibitor was performed for 2 h after plowing, MMP-3 was upregulated 7.8-fold (fig. 3b). This value was not different from the MMP-3 expression of plowed cartilage measured after 2 h of equilibration without actinomycin D (p = 0.45).

Besides MMP-3 gene regulation, the actinomycin D treatment of cartilage revealed that plowing also influences TIMP-1 expression. Upon plowing completion, TIMP-1 expression was upregulated 1.7-fold (fig. 3a); similarly, when the cartilage strips were preincubated for 2 h in media containing actinomycin D and then subjected to the 100-newton plowing, the gene expression of TIMP-1 was increased 2-fold. Moreover, the cartilage incubated in actinomycin D for 2 h after plowing induced a strong (7.5-fold) TIMP-1 upregulation (p = 0.0010; fig. 3b). Since qRT-PCR of plowed cartilage strips not
treated with inhibitor revealed that TIMP-1 expression measured after 2 h of equilibration was unaffected by plowing, these results suggest that when transcription is enabled, the TIMP-1 mRNA content is reduced. The mRNA content of Coll1 was similar to that of TIMP-1, indicating that plowing also reduces the stability of Coll1 mRNA (p = 0.0010). The other studied genes, Agg, Fn and Coll2 which, normally, were only slightly affected or not affected by plowing when no inhibitor was added, showed a different time course. Indeed, when incubation in actinomycin D was performed either before or after plowing, levels of expression of Agg, Fn and Coll2 remained at

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**Fig. 2.** Gene regulation after plowing with applied normal forces of 25, 50 and 100 N. Black bars: gene expression after 2 h of equilibration after plowing. Grey bars: gene expression after 4 h of equilibration after plowing. White bars: gene expression after 24 h of equilibration after plowing. Expression levels (normalized against the GAPDH and 18S-rRNA genes), are normalized to those of controls (i.e. nonloaded, free-swelling explants). Each experiment was from a separate bovine nasal septum (n = 3), and 3 similarly treated explants were pooled for RNA extraction in each. * p < 0.05.
around 2-fold upregulation. In particular, the statistical analysis revealed that the expression of Agg increased due to the treatment with actinomycin D (p = 0.009 and p = 0.007 for treatment before and after plowing, respectively). Similar behavior was observed for Fn (p = 0.010 and p = 0.017 for treatment before and after plowing, respectively) and Coll2 (p = 0.011 and p = 0.004 for treatment before and after plowing, respectively).

**Effect of Plowing on GAG Release**

To determine if GAG release after plowing was mechanically or enzymatically induced, MMP activity was inhibited by GM6001 treatment of the cartilage explants (i.e. controls and plowed samples) before and after plowing. As shown in figure 4, all samples showed increasing GAG release over time with excellent coefficients of determination ($R^2$ values $\geq 0.9$).

GAG release from plowed samples, independently of the GM6001 treatment, was higher than that from the corresponding controls.

In addition, the GAG release from the control samples showed no statistically significant difference, but the plowed samples treated with GM6001 or not treated showed statistically significant differences of GAG release over time (for day 1: p = 0.011, day 2: p = 0.003, day 3: p = 0.018 and day 4: p = 0.012). These results suggest that GAG release is due to both a mechanical and an enzymatic component, and that under plowing conditions, the enzymatic component is dominant.
Discussion

In this study, we performed plowing of pristine cartilage strips and demonstrated that, in vitro, this causes cell death at the surface of the explants, changes the chondrocyte gene expression and also induces GAG release by activating catabolic enzymes.

Bovine nasal cartilage was selected as model tissue because of its unique features: it is pristine (not previously subjected to any mechanical load), has homogeneously distributed chondrocytes and, along with being easily available in big quantities, it can be easily shaped. In addition, it was recently demonstrated that bovine nasal cartilage behaves as a biphasic material and has viscoelastic responses to dynamic forces [Colombo et al., 2013]. During jaw opening/closing, the TMJ cartilage disk is mainly affected by the plowing forces that are the dominant components of the tractional force. The plowing of the disk results from the combination of an applied normal force and the sliding of the condyle.

The plowing parameters used in this study were comparable to those encountered under physiological conditions. The indenter speed was chosen after evaluation of TMJ recordings performed during rhythmic jaw opening and closing [Gallo et al., 2000]. Interestingly, this speed value can also be compared to that estimated in other joints, such as in the knee during walking [Waldman and Bryant, 1997]. The applied normal forces were chosen according to the study of Sellers and Crompton [2004], showing that 100 N corresponds to the condylar TMJ force occurring during biting.

We provided here a model of cell death caused by plowing. We found that the plowing of cartilage strips induces cell death progressively, increasing along with the magnitude of applied normal force. The fact that dead cells were mainly detected at the surface of the explants suggests that chondrocytes located closer to the surface are more exposed to mechanical stress and are therefore more vulnerable than those in the deeper zones. Furthermore, qRT-PCR revealed that MMP-3 upregulation, similar to the superficial cell death observed, is dependent on the magnitude of the applied mechanical stress and that it decreases over a 24-hour equilibration period after plowing (fig. 2). In contrast, the mRNA content for Agg, TIMP-1, Fn, Coll1 and Coll2 remained unchanged compared to the control values. As active MMP-3 digests collagens, proteoglycans and other extracellular matrix proteins as well as activating the proforms of other MMPs and aggrecanase II [Murphy et al., 2002; Cawston and Wilson, 2006; Echtermeyer et al., 2009], our results suggest that plowing causes an increase in catabolic activities starting at 25 N of applied normal force.

It is well known that cartilage has a poor intrinsic healing capacity [Lima et al., 2004]. Nevertheless, after an injury, the healthy chondrocytes promote a remodeling process that consists of the elimination of the damaged matrix and the rebuilding of new matrix [Treadwell et al., 1991]. We could thus suppose that in the plowed cartilage, viable chondrocytes start to remodel the matrix by producing MMP-3, as shown by qRT-PCR experiments, in order to clear space for cell ingrowth and/or the deposition of newly synthesized proteins.

Application of the transcription inhibitor actinomycin D before plowing reduced the MMP-3 mRNA content to its control level (fig. 3a), indicating that early events during the 2-hour plowing period induce an increase in MMP-3 mRNA transcription. Under the same conditions, the mRNA of TIMP-1, Fn, Coll1 and Coll2 were increased by a factor of 2. Assuming that, in the presence of actinomycin D, the transcription was fully inhibited (as in the case of MMP-3), these results suggest that all these mRNAs were, to a certain extent, prevented from degradation during plowing and that this process is, in
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...turn, dependent on transcription. Considering that all these mRNA content data were compared to controls that did not undergo plowing, it is suggested that plowing has an additional, so far unknown, stabilizing effect on mRNAs for TIMP-1, Coll1, Agg, Fn and Coll2, that depends on active transcription. It could also be that actinomycin D itself stabilizes certain mRNA species. The degradation of mRNA is an essential determinant in the regulation of gene expression, and it can be modulated in response to environmental signals by cis-acting elements and trans-acting factors that contribute to mRNA regulation decay [Tourriere et al., 2002; Simon et al., 2006]. In addition, it has already been reported that actinomycin D has a stabilizing effect on the mRNA transcription of TIMP-1 by affecting trans-acting factors involved in TIMP-1 mRNA degradation [Gardner et al., 2006]. This stabilizing effect or artifact of actinomycin D could also account for the 2-fold increase in Agg, Coll2 and Fn mRNA when applied for 2 h upon completion of plowing.

The 7- to 8-fold increase in mRNA content for TIMP-1 and Coll1, however, suggests that both species increased during plowing and that, in the absence of actinomycin D, they decreased to control levels during the 2-hour equilibration period. This hypothesis would require a postplowing half-life of TIMP-1 and Coll1 mRNA of <1 h. In contrast, the half-life of MMP-3 mRNA is about 24 h (fig. 2). Whether or not plowing can induce factors that are able to modulate the half-life of specific mRNA species such as TIMP-1 or Coll1 cannot be definitively determined from these results.

In essence, we have observed three patterns of transcription regulation. The first is of the MMP-3 mRNA type, the transcription of which is induced during plowing. The second includes TIMP-1 and Coll1, the mRNA decay of which occurs early in the equilibration period after plowing. The third applies to Agg, Fn and Coll2, the basal transcription of which is either stabilized or unaffected by plowing.

Furthermore, we showed that the amount of GAG release from the plowed cartilage strips was higher than the corresponding release from control explants [DiMicco et al., 2004; Lin et al., 2004], and that this release was due to both mechanical damage and enzymatic activity. Since GAG release was also observed after the treatment of plowed cartilage with GM6001, we could state that plowing provokes the mechanical degradation of the cartilage matrix. This finding is also supported by two macroscopic events, namely, the bending of the strip during plowing (due to the fact that the strip is glued by the extremities to the support) and the increase in the length of the cartilage strips. The cartilage strip deformation strongly depended on the applied normal force during plowing. An increase in the entire length of the sample was observed at all the applied forces. In more detail, for samples plowed with the normal forces of 25, 50 and 100 N, a length increase was measured of 0.5 ± 0.07, 1.0 ± 0.06 and 1.9 ± 0.29 mm, respectively; this corresponded with a relative strip elongation of 0.7 ± 0.3, 1.5 ± 0.2 and 3.0 ± 0.7% for 25, 50 and 100 N, respectively.

Both observations suggest that plowing could damage the collagen fibers, causing, in turn, a GAG release [DiMicco et al., 2004]. Nevertheless, given that even after 1 day of equilibration, the GAG release from plowed cartilage was 30% higher than in the plowed cartilage treated with GM6001, we could conclude that GAG loss is also due to the catabolic activity of MMPs that are activated during plowing. Thus, the increase in MMP mRNA induced by plowing, as in the case for MMP-3, yields an increase in MMP activity.

When cartilage explants are subjected to a static compression with an applied pressure >0.5 MPa, the chondrocyte metabolism is already irreversibly compromised after 1 h [Sah et al., 1989; Valhmu et al., 1998]. The pressure during the 100-newton plowing has been calculated to be around 2.5 MPa, taken from measurements of the contact area between the condyle and the cartilage. This finding suggests that plowing with a high applied pressure is not as harmful as a static compression. In contrast to what happens during static compression where the flow of nutrients is limited, during plowing, the sliding of the indenter not only mixes the surrounding medium and thus facilitating the exchange of molecules and ensuring better cartilage homeostasis, it also squeezes fluid out of the cartilage which is subsequently replenished by new medium.

The response of cartilage explants to mechanical injuries has been extensively investigated in vitro, but, to our knowledge, plowing experiments where the indenter simultaneously applies a compression and slides on the cartilage explants have never been reported.

On the basis of the obtained results, we deduce that plowing with an applied normal force of 100 N and an indenter speed of 10 mm/s causes cell death of the chondrocytes closer to the surface as well as matrix damage, observed as GAG loss. In addition, in healthy chondrocytes, plowing promotes the production and activation of catabolic enzymes like MMP-3 and, 2 h after plowing, shows no effect on anabolic genes like Agg, Coll1, Coll2 and Fn.

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