INTRODUCTION

Articular hyaline cartilage is a specialized connective tissue important for the distribution of pressure and shear forces within joints with a specific role in animal locomotion.
Mature articular cartilage has a limited potential for the repair of critical-sized defects because of its avascularity and the absence of stem cells [2]. The response of the normal articular cartilage to damage or degeneration is frequently partial and repair of the damaged tissue is incomplete. Surface defects of articular cartilage do not heal spontaneously and usually turn to extensive degeneration [3]. It was observed that despite the limited ability of chondrocytes to migrate from marginal zones into the defective site, they cannot multiply in these locations and produce macromolecules needed for the repair of the organized intercellular matrix typical of normal articular cartilage [4].

Cartilage properties result from its microscopic structure. Detailed studies of morphology and biology of mature cartilage showed that it is a highly organized structure. Complex connections between the chondrocytes and the matrix are maintained actively and are responsible for the preservation of the shape and function of the tissue owing to the structural character of the solid component (e.g.: collagen, proteoglycans and glycoproteins). Therefore, the tissue that would reconstruct correctly the damaged surface of the articular cartilage has to replicate these structures very precisely thus cartilaginous tissue defects could be repaired.

Current scientific trends present new possibilities for the repair of cartilage defects. Transplantation of autologous or allogeneic chondrocytes, synovial mesenchymal stem cells, and biodegradable scaffolds are the methods that have been tested experimentally and clinically [5-11]. Implants seeded with chondrocytes cultured \textit{in vitro} support the production of the extracellular matrix by chondrocytes, mainly glycosaminoglycans and type II collagen [12,13]. The implantation of chondrocyte–collagen composites into cartilage defects has proved a promising method of cartilage repair [14,15] and numerous \textit{in vivo} studies in animals have shown that hyaluronan-based scaffolds seeded with autologous chondrocytes are useful for inducing the formation of hyaline-like cartilage tissue and are reabsorbed in the absence of an inflammatory response [16,17].

The aims of our study were to investigate a repair process of artificially created cartilage defects in rabbits treated by autologous and allogeneic chondrocytes in a suspension in fibrin glue Beriplast\textsuperscript{*} or seeded on Collagen type I-Hyaluronan (Col type I-HYA) scaffolds. Chondral defects (ChD’s) were created in such a manner as not to affect the zone of calcified cartilage in a non-load-bearing area of the medial condyle of the distal femur. The healing process of ChD’s was evaluated macroscopically, as well as by histological and immunohistochemical analyses 12 months after the treatment.

**MATERIALS AND METHODS**

**Animal care**

This study was performed at the University of Veterinary Medicine Košice, Slovakia respecting the guidelines for animal experiments with the approval of the Ethical Committee. This experiment was carried out on 15 domestic rabbits assigned to four treated groups and the untreated “control” group (n = 3 in each). Within four weeks, two surgical procedures were performed. In both cases, the rabbits were premedicated with an intramuscular injection of 1% ketamin hydrochloride (Narkamon inj. a.u.v.,
Léčiva, CZ) and 2% xylasin hydrochloride (Rompun amp. a.u.v, Bayer, Turkey) at doses of 40 mg/kg BW and 4 mg/kg BW, respectively and anesthesia was maintained with 2.0 % isoflurane at 2 l/min. constant oxygen flow.

The first surgery was performed on the knee joint of the left hind leg to obtain articular cartilage from a non-weight bearing area of the medial femur condyle. All samples of articular cartilage were harvested for 4 weeks in vitro cultivation of chondrocytes. One month after the first surgery, on the same knee joint, circular chondral defects (3 mm in diameter) were created artificially at the non-weight bearing area of the medial femur condyle using a metal puncture needle (Chirana, SR) during arthrotomy of the knee joint. The ChD’s were made of a desirable size and shape. While creating the ChD’s, our aim was to prevent their penetration into the vascularized subchondral zone as the zone of calcified cartilage should remain intact, because deformations of the vascularized layer are accompanied by a bloody discharge what induces an “undesired” non-specific healing process.

One month after the first surgical procedure, the second operation was performed under identical conditions, using the same procedure as described previously. The ChD’s created artificially during the first surgery were treated as follows:

1st group: The ChD’s were filled with autologous chondrocytes seeded on Col type I-HYA scaffolds. The scaffolds were fixed to the ChD’s by fibrin glue Beriplast® (Centeon Pharma GmbH). The scaffolds corresponded to the size, shape and depth of the chondral defects, and they were implanted into the ChD’s in such a fashion that the surface seeded with cultured chondrocytes was directed into the defect base, while the surface without cultured chondrocytes was oriented to the synovial cavity of the femur.

2nd group: The ChD’s were filled with allogenous chondrocytes seeded on Col type I-HYA scaffolds. The same way as in Group 1 were used for implantation of the scaffolds.

3rd group: The ChD’s were filled with a suspension of autologous chondrocytes in fibrin glue Beriplast® and they were immediately covered by Col type I-HYA scaffolds.

4th group: The ChD’s were filled with a suspension of allogenous chondrocytes in fibrin glue Beriplast® and they were immediately covered by Col type I-HYA scaffolds.

Control group: The ChD’s were left to heal spontaneously without any treatment. Immediately after treatment, the knee joint was stabilized with a modified Robert Jones bandage for 2 weeks. All rabbits were sacrificed by an intravenous injection of veterinary euthanasia solution (T-61, Hoechst, Somerville, NJ) for the collection of samples 12 months after the treatment.

Allogenous and autologous chondrocyte isolation and culture, and Col type I-HYA Scaffold preparation

Specific procedures for the isolation and cultivation of chondrocytes and the preparation of Col type I-HYA scaffolds were as described by Harvanova et al. [18]. Briefly, circular fragments of rabbit cartilage (3 mm in diameter) were digested by bacterial collagenase
type II (1mg/mL) (Gibco BRL®) in Nutrient mixture F-12 medium (Gibco BRL®) supplemented with 1% (v/v) antibiotic/antimycotic solution for 16-20 hours. The obtained chondrocytes were collected and cultivated in Nutrient mixture F-12 medium supplemented with 20% (v/v) fetal bovine serum (Gibco BRL®), 1% (v/v) insulin-selenium-transferin A (Gibco BRL®), 1% (v/v) antibiotic/antimycotic solution at 37°C in humidified air with 5% CO₂ for one month. Isolated chondrocyte cultures were passaged before confluence was obtained with trypsin-EDTA (Gibco BRL®) and the medium was exchanged every three days. In the 3rd and 4th group, autologous and allogenous chondrocytes were used for the *in vivo* application with fibrin glue Beriplast® at density of 0.65-0.8 x 10⁶ per ml and seeded on the Col type I-HYA scaffold, respectively.

Enzymatically treated fibrous collagen type I (bovine Collagen type I from Achilles tendon, Hypro, Otrokovice, Czech Republic) was mixed with 0.5 M solution of acetic acid and aqueous solution of sodium hyaluronate (10mg/ml, 1500 kDa, CONTIPRO Ltd., Dolní Dobrouč, Czech Republic). Starch dialdehyde 2% (v/v) aqueous solution was added to the collagen/hyaluronan complex and homogenized in a blender at 1400 RPM for 1 minute, pre-frozen for 12 hours and freeze dried (JOUAN LP3) at -45°C, 2x10⁻² mBa for 6-8 hours. After lyophilisation, scaffolds were sealed in polyethylene bags and sterilized by ⁶⁰Co radiation with a dose of 30 kGy. Lyophilized scaffolds were cut to a desired shape (3 mm in diameter) and were left in the culture medium for 1 hour before *in vitro* cultivation. In the 1st and 2nd group, autologous and allogenous chondrocytes were seeded on the Col type I-HYA scaffold in at density of 0.47-0.6 x 10⁶ per cm² and were cultured at 37°C in humidified air with 5% CO₂ for 4 weeks, respectively. The culture medium was exchanged every three days.

**Locomotory activities**

Evaluation was based on the time needed to full load limb recovery after treatment and on monitoring of patella position, septic arthritis, production of exudate compactness of suture, and infection of wounds which can also reflect the status of the defect repair.

**Macroscopic evaluation**

The gross appearance of the ChD’s was evaluated for color, integrity, contour and smoothness of the surface.

**Histological and immunohistochemical analyses**

The specimens of treated medial condyles of the distal femur in each rabbit were fixed in 10 % phosphate-buffered formalin (Sigma-Aldrich®, Slovak Republic) for 48 h. Subsequently, they were washed in water and decalcified in 25% Chelaton III (v/v) pH 7.4 (Bioptika, Italy) at 37°C for 7 days. After decalcification, the samples were washed in water for 24 h, embedded in paraffin wax, and 15 μm thick slices were cut sagittally, and were stained *in situ* with Haemotoxylin-Eosin (H&E) and Safranine O (specific for proteoglycans). Immunohistochemical staining was carried out using antibodies against collagen type II fibres (Rabbit Anti-Human Collagen type II, BIODESIGN
International, Saco, ME USA) and porcine immunoglobulin against human IgG tagged with fluorescein-isothiocyanate (SwAHu/IgG-FITC, SeVac, Prague, Czech Republic). Both antibodies were diluted 1:100 in 1% phosphate buffered saline (10 mM sodium phosphate, 0.15 M sodium chloride, pH 7.4). The evaluation was performed by light microscopy and fluorescence microscopy using Olympus advanced inverted microscope IX-71 with an improved fluorescence illuminator. The histological sections were examined for the quality of the repair process and were graded with use of histological scale described by [6]. All stained slides were evaluated in a blind test by three experienced histologists.

RESULTS

Locomotory activities
Despite repeated arthrotomy, none of the animals showed complications in relation to patella luxation, septic arthritis and excessive production of exudate, dehiscence of sutures or secondary infection of wounds. After the second operation, we observed that the rabbits spontaneously put less weight on the operated limb. The full load and extension of the hind leg recovered completely at the end of the 3rd week after treatment, it means, one week after the bandage removal. Moreover, pain as a factor which significantly limits movements, is not expected in this type of defect (partial cartilage loss) due to the avascular and aneural character of the articular cartilage.

Macroscopic evaluation
The patellar joints were filled with a clear viscous synovial fluid in all rabbits. In all treated groups, neither detachment of the Col type I-HYA scaffolds (except for a severally irregular surface of the treated ChD in one rabbit in the 4th group) nor signs of articular surface damage (such as chondromalacia, osteoarthritis or synovitis) or hypertrophy were found. In the 1st and 3rd group treated by the autologous or allogenous chondrocytes, respectively, the ChD´s have been filled with smooth cartilaginous tissue of a color similar to healthy cartilage (Fig. 1a). The ChD´s were completely filled in 3 rabbits (one in the 1st group and two in the 3rd group), but one edge of the scaffolds was not completely integrated with the ChD´s in the other rabbits. On the other hand, in the 2nd and 4th group treated with the suspension of autologous or allogenous chondrocytes, respectively, we have found just partial healing of the ChD´s in all rabbits (Fig. 1b). In the control group, no signs of cartilage reparation processes were observed and the ChD´s retained sharp edges even 12 months after surgery (Fig. 1c). Surprisingly, in the rabbit no. 15, the ChD was covered with a whitish tissue that did not resemble the cartilage tissue.

Histological and immunohistochemical analyses
Results of the histological examination of all stained slides are presented in Table 1. In the 1st group, the ChD´s were filled with a hyaline-like cartilage tissue (HyLCa) (Fig. 2a), but the intensity of Safranin O staining was high in all layers (deep, middle and
surface layer) of the ChD (Fig. 2b) only in rabbit no. 3. Except for HyLCa, we have also observed sporadic segments of fibrocartilage (FiCa), particularly in the surface layer of the ChD’s in rabbit no. 1 and no. 2 from the same group. Safranin O staining revealed a relatively high production of proteoglycans (PG’s) and collagen type II fibres, in the deep layer, but lower intensity of staining has been exhibited in the surface layer of the ChD’s. Chondrocytes in the deep layer have a round shape and their density was higher as compared to the surface layer of the ChD’s. Immunohistochemical staining confirmed the presence of cells with the chondrocyte phenotype in all ChD’s (Fig. 2c).

In rabbit no. 1, a slight hypertrophy of HyLCa was detected probably due to the shape of the scaffold which could be not completely suitable for the insertion into the defect. All in all, a local inflammatory reaction was not observed in all cases.

In the 2nd group, apparently lower numbers of cells with the chondrocyte phenotype occupied the ChD’s (Fig. 3a), as well as decreased production of PG’s (Fig. 3b) and collagen type II fibers (Fig. 3c) were found than in the 1st group. In addition, the ChD’s healed with both types of cartilage tissues, but predominantly by FiCa. A local

Figure 2. Histological and immunohistochemical analyses of the chondral defects repaired by autologous chondrocyte seeded on Col type I-HYA scaffold (1st group) a) H&E staining, 150x magnification b) Safranin O staining, 100x magnification, arrow shows a weakened adhesion of the autologous chondrocyte seeded on Col type I-HYA scaffold to the native chondral cartilage c) Immunohistochemical staining, 200x magnification. (HyLCa = hyaline-like cartilage tissue, 1—native chondral cartilage, 2—autologous chondrocytes seeded on Col type I-HYA scaffold)

Figure 1. Macroscopic view of the chondral defects repaired a) by autologous chondrocyte seeded on Col type I-HYA scaffold (1st group), b) by suspension of allogenous chondrocytes in a fibrin glue (4th group) c) without any treatment (Control group). Arrows show the position of the chondral defects.
inflammatory reaction was not confirmed by histological analysis, what indicates that the allogenous chondrocytes did not cause the scaffold rejection accompanied by its lacked mechanical stability in the defect leading to inflammation.

In the 3rd group, the ChD’s were filled completely with HyLCa in two rabbits (no. 8 and no. 9). In the deep layer of the ChD’s, the cells resembling chondrocytes of the native cartilage tissue have a round shape (Fig. 4a, 4b). The surface layer of the ChD’s showed the presence of collagen type II fibers produced by chondrocytes seeded on Col type I-HYA scaffold as in the 1st group. The intensity of PG’s was low in the surface layer, particularly at the edges of the ChD’s, but in the middle and deep layer were higher (Fig. 4c, 4d). Immunohistochemical staining proved the presence of a cellular population of the chondrocyte phenotype in the ChD’s (Fig. 4e).

In the 4th group, we have observed only partial regeneration of the ChD’s in two rabbits (no. 10 and no. 12). A severely irregular surface of the treated ChD was found in the rabbit no. 11. Reparation of the ChD’s was incomplete and damage was located mostly at the edges of the defects. HyLCa has been observed in the middle and deep layer of the defects, but the periphery of the ChD’s was filled with FiCa. The production of PG’s and the density of cells with the chondrocyte phenotype were low. Fibrin glue components without the cell population were seen in these ChD’s, too.

Table 1. Histological grading scale for the repair of the chondral defects in individual rabbits.

<table>
<thead>
<tr>
<th>Category</th>
<th>Points</th>
<th>1st group</th>
<th>2nd group</th>
<th>3rd group</th>
<th>4th group</th>
<th>Control group</th>
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<tr>
<td>Number of rabbit</td>
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<td>1 2 3</td>
<td>4 5 6</td>
<td>7 8 9</td>
<td>10 11 12 13 14 15</td>
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<td>Hyaline cartilage</td>
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<td>Mostly hyaline cartilage</td>
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<tr>
<td>Mostly fibrocartilage</td>
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<td>Mostly non-cartilage</td>
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<td>Non-cartilage</td>
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<td>Slightly reduced</td>
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<td>Markedly reduced</td>
<td>2 2 2 2</td>
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<td>No metachromatic staining</td>
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<td>Surface regularity</td>
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<td>Smooth (&gt;3/4)</td>
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<td>Moderate (&gt;1/2-3/4)</td>
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<td>Irregular (1/4-1/2)</td>
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<td>Severely irregular</td>
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<td>Thickness of cartilage</td>
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<tr>
<td>&gt;2/3</td>
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<td>1/3-2/3</td>
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<td>&lt;1/3</td>
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<td>Integration of donor with host adjacent cartilage</td>
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<td>Both edges integrated</td>
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<tr>
<td>One edge integrated</td>
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<tr>
<td>Neither edge integrated</td>
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<td>8 8 7 4 0</td>
<td>8 9 8 14 14 10</td>
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</table>
In the control group, the ChD’s were left to heal spontaneously. The defects were empty in 2 rabbits (no. 13 and no. 14). In the rabbit no. 15, non-specific repair mechanisms were observed in the deep layer of the ChD by the formation of FiCa. Neither the production of PG’s nor collagen type II fibers were found in this group.

**Figure 3.** Histological and immunohistochemical analyses of the chondral defects repaired by allogeneous chondrocyte seeded on Col type I-HYA scaffold (2nd group) a) H&E staining, 150x magnification, b) Safranin O staining, 100x magnification, very low production of PG’s. c) Immunohistochemical staining, 100x magnification, production of collagen type II localised mainly in deep layer of the chondral defect. (PG’s = proteoglycans, 1-native chondral cartilage, 2-allogeneous chondrocytes seeded on Col type I-HYA scaffold, 3 – subchondral bone)

**Figure 4.** Histological and immunohistochemical analyses of the chondral defects repaired by a suspension of autologous chondrocytes in fibrin glue (3rd group) a) H&E staining, 100x magnification b) H&E staining, 150x magnification, c) Safranin O stain, magnification 100x, surface layer created by collagen type I fibres from the scaffold, and marginal zone of the chondral defect with a low production of PG’s, and deep layer of the defect is repaired by HyLCa d) Safranin O stain, magnification 150x, deep and middle layer of the chondral defect with high production of PG’s d) Immunohistochemical staining, 150x magnification. (HyLCa = hyaline-like cartilage tissue, PG’s = proteoglycans, 1 – surface layer of the chondral defect, 2 – deep layer of the conndral defect, 3 – deep layer of the native chondral cartilage, 4 – subchondral bone, arrows indicate the marginal zones of the chondal defect)
DISCUSSION

In the last few years, a great effort was made to define a suitable method for the implantation of chondrocytes into the defective cartilage in order to support the regeneration of cartilage tissue. Studies comparing methods for cartilage tissue reconstruction using autologous chondrocytes, as well as allogeneic chondrocytes are lacking. Therefore, we have decided to use two different methods of implantation of autologous and allogeneic chondrocytes into the ChD’s: (a) a suspension of chondrocytes in a fibrin glue covered by the Col type I-HYA scaffolds, and (b) chondrocytes seeded on the Col type I-HYA scaffolds fixed with a fibrin glue. In addition, Lee et al. [19] found that chondrocytes cultured for 4 weeks are able to produce higher amounts of hyaline cartilage after their implantation into chondral defects than these cultured for a significantly shorter time (only 12h). Therefore, the culturing period of chondrocytes for one month was chosen in our study.

Different scaffolds were used as cell or tissue carriers for cartilage tissue engineering so far [20-23]. The major advantage of collagen scaffolds, which were used in our experiment, is their tissue abundance and a highly organized structure. It is known, that porous three-dimensional structures consisting of collagen fibers support the viability of chondrocytes and their production of extracellular matrix in vitro. Collagen contains cell-adhesion domain sequences that induce specific cellular interactions [24]. Substrate-bonded hyaluronan exhibits size-dependent stimulation of chondrogenic differentiation and also improves the proliferative activity of cultured chondrocytes [25]. As previously published [26-28], small amounts of hyaluronan (2% w/w) in three-dimensional collagen scaffolds can enhance chondrogenesis. Based on our previous study [18], we can corroborate that the Col type I-HYA scaffolds had not any adverse effects on chondrocyte viability, morphology and phenotype during long term (2-3 months) in vitro cultivation. Moreover, Lin et al. [23] found that chondrocytes seeded on Hyaluronan/Collagen II microspheres could exhibit a high activity in repairing a cartilage defect, what is in accordance with our findings.

On the other hand, suspension with chondrocytes can be injected with ease to the defect or target site. Defects orientated downwards are influenced by the gravitational forces which can affect the initial distribution of transplanted chondrocytes and may have an important role in the final clinical outcome [29]. This phenomenon was not observed in our study, but the partial detachment of the Col type I-HYA scaffold was found in one rabbit treated with the suspension of allogeneic chondrocytes in fibrin glue. By our opinion, adhesive properties of the fibrin glue could be altered due to its dilution with a medium of cultured chondrocytes. All in all, comparing autologous vs. allogeneic chondrocyte application, better proliferation of cells with the chondrocyte phenotype and production of PG’s, as well as the creation of HyLCa were observed in both groups treated with autologous chondrocytes. Hong et al. [30] hypothesized that autologous structures have better biocompatibility with native tissues, which was corroborated with our findings too.

As a limitation of this study can be considered that, a small number of experimental animals (only 3 rabbits per group) was used. This discouraged the use of statistical
analysis within the groups thus individual examination was chosen to present our findings instead. Moreover, rabbits are not an optimal animal model for the evaluation of the ChD’s due to the relatively small dimensions of the knee joint and a thin layer of chondrium, which is not thicker than 1 mm.

In all treated groups, the chondrocytes were capable to proliferate and produce the cartilage extracellular matrix, including proteoglycans and type II collagen fibers, as compared to the control “untreated” group. The fibrin glue ensured a sufficient fixation of the scaffolds when it was applied undiluted, but its fixative properties could be altered by a medium of chondrocytes added thus further investigations with a higher number of animals involved are required. For this reason, it can be concluded that the autologous chondrocytes seeded on a Collagen type I-Hyaluronan scaffold represents a very successful way of treatment of articular cartilage defects in rabbits.

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REFERENCES


27. Chang CH, Kuo TF, Lin CC, Chou CH, Chen KH, Lin FH, Liu HC: Tissue engineering-


REPARACIJA DEFEKATA HRSKAVICE KUNIĆA AUTOLOGNIM I ALOGENIM HONDROCITIMA ZASEJANIM NA KOLAGEN/ HIJALURONAN OSNOVU Ili SUSPENDOVANIM U FIBRINSKOM LEPKU

HORŇÁK Slavomír, HARVANOVÁ Denisa, LEDECKÝ Valent, HLUCHÝ Marian, VALENČÁKOVÁ-AGYAGOSOVÁ Alexandra, AMRÍCHOVÁ Judita, ROSOCHA Ján², VAŠKO Gabriel, ŠVIHLA Róbert, PETROVIČ Vladimír

Cilj studije je bio da se verifikuje in vivo preživljanje in vitro kultivisanih autolognih i alogenih hondrocita suspendovanih u fibrinskom Beriplast® lepaku ili na kolagen Hylauronan tip I (Col type I-HYA) osnovu radi regeneracije oštećenja zglobove hrskavice kod kunića. Ispitivanje je obavljeno na 15 domaćih kunića, odabranih slučajnim izborom koji su podeljeni u pet grupa (3 po grupi) kod kojih su veštački izazvana oštećenja hrskavice (ChD), tretirana na različite načine. Ostećenja su napravljena na površinama koje nisu noseće na medijalnom kondilusu distalnog femura, a tretman je podrazumijeva: prva i treća grupa: ChD su punjene sa autolognim ili alogenim hondrocitima zasejanim na Col tip I-HYA osnovu. Osnove su fiksirane na ChD pomoću fibrinskog lepka Beriplast®; druga i četvrta grupa: ostećenja su punjena suspenzijom autolognih ili alogenih hondrocita u fibrinskom Beriplast® lepku, a neposredno posle toga, prekriveni su sa nezasejanom Col tip I-HYA osnovom; kontrolna grupa: ostećenja su ostavljena da spontano zarastu, bez tretmana. Dvanaest meseci posle tretmana obavljene su makroskopske, histološke i imunohistološke analize promena. Kod svih životinja tretiranih grupa, hondrociti su bili sposobni da proliferišu i proizvode ekstracelularni matriks hrskavice, uključujući proteoglikane i II tip kolagena u poredenju sa kontrolnom “netretiranom” grupom. Sa druge strane, proizvodnja hrskavičavog tkiva nalik na hijalin potvrđuje da se oba terapijska metoda koji koriste autologne hondrocite mogu da uspešno primene u tretiranju defekata hrskavice kod kunića.