Pentosan Polysulfate as a Disease Modifier of Cartilage Degeneration in Experimental Osteoarthritis

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Abstract

Reasons for performing study: Disease modifying drugs for Osteoarthritis (OA) is an important trend of development of an appropriate therapeutic protocol for both human and animal. To date no ideal OA treatment protocol has been described.

Objectives: To assess clinical, histologic, and biochemical effects of sodium pentosan polysulfate (NaPPS) administered intramuscularly (IM) for treatment of experimentally induced osteoarthritis in donkeys.

Study design: OA was induced using repeated injection of allogenous donkey cartilage. Six donkeys received NaPPS (3 mg/kg, IM) on study days 70, 77, 84, and 91. Six control donkeys received the same volume of saline (0.9% w/v NaCl) IM on study days 70, 77, 84, and 91. Clinical, gross, histologic, histochemical, and biochemical findings as well as synovial fluid analysis were evaluated.

Methods of testing hypotheses: Lameness score, carpal circumference, joint flexion angle, Synovial fluid analysis, light and scanning electron microscopy were evaluated.

Results: Induction of osteoarthritis caused increase in the lameness score, joint circumference, synovial fluid calcium, phosphorus, magnesium, total protein and leukocyte count and cartilage damage. NaPPS treatment significantly reduced synovial fluid Ca, Mg, P and cartilage damage.

Conclusions: NaPPS resulted in significant improvement in clinical signs and articular cartilage healing, and no adverse effects were detected confirming that NaPPS has disease-modifying properties.

Potential relevance: NaPPS is a suitable therapeutic option for osteoarthritis in equines.

Keywords: Pentosan polysulfate; Osteoarthritis; Cartilage; Synovial fluid; Joint crystals; ACP model

Introduction

Equine osteoarthritis (OA) may be considered as a group of disorders characterized by a common end stage: progressive deterioration of the articular cartilage accompanied by changes in the bone and soft tissues of the joint. The deterioration of the articular cartilage is characterized by local splitting and fragmentation (fibrillation) of articular cartilage. Synovitis and joint effusion are often associated with the disease, and, clinically, the disease is characterized by pain and dysfunction of the affected joint [1]. Several epidemiologic studies have found that lameness due to joint disease is the most significant factor responsible for inability to race and loss of performance [2-4].

Drugs used to treat osteoarthritis can be considered as modifying the clinical signs of osteoarthritis or as modifying the disease itself. In general, the former alleviate pain and inflammation and lead to a positive clinical outcome but may not affect the progression of the underlying disease process. The latter slow or reverse the disease process but may not have any direct effect on the clinical signs of the disease. The optimal medication for osteoarthritis would be a drug that provides both effects in a safe and convenient manner.

Various medications have been evaluated or used for treatment of horses with osteoarthritis, including nonsteroidal anti-inflammatory drugs (NSAIDs), corticosteroids, polysulfates glycosaminoglycans (PSGAG), and hyaluronan [5].

Although NaPPS has been used in Europe for more than 30 years as an antithrombotic / antilipidemic agent, its potential as a disease-modifying antiarthritic agent has been realized more recently [6,7]. The backbone of PPS, which consists of repeating units of (1-4)-linked β-D-xylano-pyranoses, is isolated from beech-wood hemicellulose.

The possible mechanism of action of PPS are improving of the nutritional supply to the joint, increasing of biosynthesis of extracellular matrix, inhibiting of proteinases activity, enhancing TIMP-3 level in cartilage and inhibiting of inflammatory activities of IL-1 and TNFα [6].

There are numbers of clinical trials reported the use of PPS in dog [8-12], sheep [13] and human [14-16].

The pharmacology of PPS has been described in the horse [17,18] and its use for equine joint disease has been reported [19], but results

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from the later study needs further investigation in another controlled animal model.

The allogogenous cartilage particle (ACP) model was used to induce osteoarthritis in dogs [20], rabbits [21], horses [22] and donkeys [23]. Its benefits has been described recently as an in vivo model for crystal associated osteoarthritis [24].

The purpose of the study reported here was to evaluate the effects of NaPPS, administered IM, in experimentally-induced osteoarthritis by evaluation of clinical (joint lameness, range of motion, response to flexion, and synovial effusion), radiographic, gross, macroscopic, histologic, and biochemical outcome measures. Our hypothesis was that the outcome of donkeys treated with PPS would be more favorable than that of control donkeys.

Material and Methods

The experiment was approved by the Committee on Animal Experimentation at the Kafelsheikh University, Egypt (Protocol no. VET1789 - 1/4/2013).

Donkeys

The present study was performed with 13 healthy Egyptian local breed male donkeys weighting from 150 to 200 kg. Animals were housed in indoor stalls and fed on a maintenance ration of mixed grain with hay and unlimited water. All donkeys were dewormed with ivermectine 200 mcg/kg body weight (Eqvalan; Merial Limited. USA)

Prior to inclusion in the study, lameness examination, body condition, radiographs of carpal joints, range of motion of carpal joints (angle of flexion) and evidence of joint effusion were assessed to ensure that all previous variables were within normal limits (baseline measurement).

Donkeys were allowed to acclimatize for 2 weeks prior to the study. During the acclimatization period, the donkeys were trained daily to familiarize them to the experimental conditions (investigators, environment, handling and vein puncture).

Cartilage powder preparation

One local breed donkey weighting ± 150 kg was euthanized, and the articular cartilage was removed from the shoulder, carpal, fetlock, pastern, hock and stifle joints in a biosafety cabinet under aseptic conditions. The pooled cartilage was powdered under liquid nitrogen in a mortar, producing particles as small as 20 mm in diameter (able to pass easy through a 14-gauge needle). These particles were resuspended at a concentration of 50 mg/ml in a physiological saline solution contained amikacin sulfate (50 mg/ml) (Amikin; Bristol-Myers Squibb). An aliquot of the suspension was cultured for 72 hour to confirm sterility. The cartilage stock solution was stored on a fridge (4ºC) until use [23,24].

Outcome measures

Clinical examination: Clinical examinations of both right and left forelimbs were performed weekly from day 0 (baseline) throughout the study period. A professor of large animal surgery (Lameness specialist) assessed all clinical outcome variables blinded to the treatment assignment.

Lameness score: The right and left forelimb of the 12 donkeys were evaluated for lameness score on a scale 0 to 5 according to American Association of Equine Practitioners (AAEP) grading system: 0: Lameness not perceptible under any circumstances, 1: Lameness is difficult to observe and is not consistently apparent, regardless of circumstances, 2:Lameness is consistently apparent under certain circumstances, 3: Lameness is consistently observable at a trot under all circumstances, 4: Lameness is obvious at a walk, 5:Lameness produces minimal weight bearing in motion) [25].

Circumference of the carpal joint: Measurements obtained at the proximal aspect of the carpus by using of a measurement tape (in cm),

Donkeys were trotted for 15 min/day on soft ground 5 days each week until the begin of the treatment.

At day 70, the 12 donkeys were divided in to two equal groups of six. Group A (PPS group) received NaPPS 3 mg/kg (Cartrophen Equine Forte; Biopharm Australia), IM, 4 doses at days 70, 77, 84 and 91. Group B received the same volume of saline (0.9% Na Cl) IM at the same time points (Control group).

All animals had box rest for 2 weeks then had another 2 weeks of of being walked by lead (20 min daily). All donkeys were euthanized at day 119 (Figure 1).

Study design

At day 0, the remaining 12 donkeys were sedated with Xylazine HCl in a dose of 1 mg/kg (Rompun; Bayer animal health). The skin was aseptically prepared for arthrocentesis of each right radiocarpal joint to obtain synovial sample for baseline analysis. All donkeys were received 100 mg (2 mL), intra-articularly into the right radiocarpal joint using a 14G needle. These injections were repeated at 7,14,21,28,35,42 and 56 days. The left radiocarpal joint were injected with the same solution without cartilage particles (sham injection) and served as non-arthritis controls. Donkeys were trotted for 15 min/day on soft ground 5 days each week until the begin of the treatment.

Figure 1: Flowchart of the study protocol.
and with the aid of the anatomical reference points (accessory, radial, ulnar and intermediate carpal bones). Circumference was obtained weekly; hair over the selected area was clipped on a regularly scheduled basis [26].

**Maximum carpal flexion angle:** Maximum carpal flexion was measured weekly by slowly flexing the carpus until the donkey resisted. The angle was then measured in degrees with Goniometer [26].

**Synovial fluid analysis:** Synovial fluid sample (1 to 2 mL) was aseptically aspirated from each joint before each injection. The conventional analysis of synovial fluid included assessment of total protein, Calcium, Phosphorus, Magnesium concentration and white blood cell (WBC) count [27]. Total protein concentrations, Calcium, Phosphorous and Magnesium were determined via Double Beam UV Visible Spectrophotometer and commercial kits however, the WBC was counted using an automated cell counter. A clinical pathologist who was blinded to the treatment protocol did the synovial analyses [24,27].

**Gross pathology:** Animals were euthanized by administration of pentobarbitone sodium (100 mg/kg IV) (Lundbeck Inc.). Both articular surfaces were photographed, directly perpendicularly, from a standard distance of 10 cm with a digital camera (Sony Cyber-shot DSC-W350) and the photographs were edited with a photo editing software (Adobe Photoshop CS2) in order to outline both the osteoarthritic lesions and the total articular surface. Tissue samples collected from joint capsule and articular cartilage (Figure 1). The right and left carpal joints specifically examined for degree and location of articular cartilage fibrillation or erosion. A subjective grade (scale of 0 to 4) assigned for partial- and full-thickness cartilage erosion as well as synovial membrane hemorrhage. A total erosion score assigned, also with a scale of 0 to 4. For each of the 2 variables, grade 0 represented no pathological change and 4 represented a severe change. Each joint also assessed for the presence of synovial adhesions [28].

**Light microscopy:** Specimens from the synovial membrane and joint capsule harvested and placed in neutral-buffered 10% formalin (NBF) stained with H&E and examined microscopically. Samples evaluated for cellular infiltration, synovial intimal hyperplasia, subintimal edema, subintimal fibrosis and subintimal vascularity. Each variable was graded and reported as a numeric value 0 to 4 (0 = normal, 1 = slight change, 2 = mild change, 3 = moderate change, and 4 = severe change [28].

Full thickness articular cartilage samples of 5-mm3 diameter were obtained from each joint (Figure 2). Sampling sites were chosen to represent an area of dorsal cartilage (C1 and C3) and palmar cartilage (C2 and C4). Samples were placed in (NBF) for 7 days then in 10% (w/v) EDTA for 21 days for decalcification then processed routinely to a standard distance of 10 cm with a digital camera (Sony Cyber-shot DSC-W350) and the photographs were edited with a photo editing software (Adobe Photoshop CS2) in order to outline both the osteoarthritic lesions and the total articular surface. Tissue samples collected from joint capsule and articular cartilage (Figure 1). The right and left carpal joints specifically examined for degree and location of articular cartilage fibrillation or erosion. A subjective grade (scale of 0 to 4) assigned for partial- and full-thickness cartilage erosion as well as synovial membrane hemorrhage. A total erosion score assigned, also with a scale of 0 to 4. For each of the 2 variables, grade 0 represented no pathological change and 4 represented a severe change. Each joint also assessed for the presence of synovial adhesions [28].

**Scanning electron microscopy:** Cartilage C2 and C4 pieces obtained from the distal redial surface of each right carpal joint were fixed in 2.5% (v/v) glutaraldehyde solution in phosphate buffer saline (PBS pH 7.4) for approximately 24 h, followed by thorough washing using multiple changes of PBS. Secondary fixation was in a solution of 1% (w/v) osmium tetroxide in PBS for 1.5 h. Specimens were again rinsed in multiple changes of PBS then dehydrated with an ascending ethanol series 50%, 60%, 70%, 80%, 90%, 100%, followed by pure, dry acetone. This was followed immediately by critical point drying with liquid carbon dioxide, coating with gold and examination by a professor of pathology using a JEOL JSM 5200LV electron microscope. Cartilage surface was examined for erosion and micro cracks [30] and synovial membrane was examined for synovial villi destruction. Variable was graded and reported as a numeric value 0 to 4 (0 = normal, 1 = slight change, 2 = mild change, 3 = moderate change, and 4 = severe change [24].

**Statistical analysis:** Variables including carpal flexion angle, carpal circumference, TP and TWBC analyzed using a repeated measures analysis of variance (ANOVA) model with IBM SPSS (Version 23). Any test with a P value < 0.05 was declared statistically significant. When individual comparisons were made, Bonferroni post hoc test was used and P < 0.05 was considered significant. The Kruskal-Wallis non-parametric ANOVA was used to evaluate statistical differences in lameness score, gross pathological and histo-pathologic scores. Values are reported as mean ± standard deviations.

**Results**

**Clinical examination**

**Lameness score:** There was an increase in the lameness scores of the right forelimb in all donkeys at day 70 (1.67 ± 0.47). In PPS group, lameness begin to decrease from day 77 (1 ± 0) till return to the base line at day 105. However in control group, the score begin to decrease at day 112 (1 ± 0) but did not return to its baseline analysis (Figure 3A). There was a significant difference between the right forelimb of control and PPS-treated groups at study day 77, 84, 91, 98 and 105 (P < 0.001).

**Carpal circumference:** Carpal circumference was increased in right forelimb of all donkeys at day 70 (24.2 ± 0.5). Then begin to decrease at day 77 in PPS Group (23.7 ± 0.7) and at day 98 for control group (24.3 ± 0.7). At the end of the study, the circumference did not return to its baseline in right forelimb of both groups (Figure 3B). There was no significant differences between the two groups at the end of the study (P=0.126).

**Maximum carpal flexion angle:** The flexion angle decreased in the right forelimb of both group at day 70 (150 ± 4) for PPS group and (148 ± 3) for control group. The angle began to increase in day 77 in PPS group (155 ± 4) and in day 91 in control group (149 ± 3). At the end of the study the angle return to its baseline analysis in PPS group and control group (Figure 3C). There was a significant difference between PPS and control group at study day 77, 84, 91, 98 and 105 (P<0.05).
Synovial fluid analysis: Synovial WBCs count was increased in PPS group (400 ± 81 cells/µL) and control group (433 ± 74 cells/µL) at day 70 throughout the study but with no significant differences between the two groups.

Total protein also increased in PPS group (3 ± 0.1 g/dL) and control group (2.9 ± 0.1 g/dL), at day 70 throughout the study but with no significant differences between the two groups.

Calcium (Ca) level was significantly increased from day 35 in PPS group (2.7 ± 0.4) and control group (2. ± 0.2) reaching its maximum level at day 70 in PPS group (3 ± 0.3) and control group (3.3 ± 0.3). Calcium level begin to decreased at day 77 (2.7 ± 0.2) till the end of the study. There was a significant difference between the right forelimb of both group from day 77 to the end of the study (P<0.05) (Figure 4C).

Gross pathology: The right joints of both groups had high cartilage erosion score (PPS group 3.33 ± 0.47; control group 4 ± 0) compared to the left one (0 ± 0). In PPS group most of the lesions are limited to C3 region however C1 and C3 characterized by fibrous like membrane (Figure 5A). In control group partial and full thickness cartilage erosion distributed all over the joint region (Figures 5B and 5C). There was no significant difference between the two groups (P=0.138). Synovial membrane haemorrhage improved in right PPS joints (0.67 ± 0.47) compared to the right control (2.67 ± 0.47). There was a significant difference between the right joints in both groups (P<0.05). Synovial fluid calcium over the study period (Figure 4A). Phosphorus (P) level was significantly increased from at day 28 in PPS group (10.9 ± 0.7) then begin to decrease at day 77 (11.7 ± 0.7) till the end of the study. There was a significant difference between the right forelimb of both groups from day 77 to the end of the study (P<0.05) (Figure 4B). Magnesium (mg) level was significantly increased at day 21 in PPS group (2.3 ± 0.3) and at day 14 in control group (2.5 ± 0.4) reaching its maximum level at day 70 in PPS group (3.9 ± 0.3) then begin to decrease at day 77 (3.8 ± 0.3) till the end of the study (2.3 ± 0.4). However, in the control group magnesium continued to increase to the end of the study (6.5 ± 0.6). There was a significant difference between the right forelimb of both group from day 77 to the end of the study (P<0.05) (Figure 4C).

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Light microscopy of synovial membrane: There was no significant difference in synovial membrane intimal hyperplasia, or subintimal edema between right and left joints of both groups (P=1.000).

Synovial membrane cellular infiltration was increased in right control joints (3.33 ± 0.47) (Figures 6A-6F) and right PPS treated joints (1.33 ± 0.47) both groups were significantly different (P=0.003). The right control joints characterized by large cartilage particles embedded inside the subintimal layer surrounded by a severe zone of cellular infiltration (Figures 6E and 6F). Right PPS joints had mild zone of cellular infiltration around some cartilage particles (Figures 6A-6C).

Synovial membrane vascularity was increased in right PPS joints (1.67 ± 0.47) (Figures 6A and 6B), and right control joints (3.50 ± 0.50) (Figure 6E) compared to left control joints (P < 0.001). Right PPS and control joint were statistically different (P=0.012).

Subintimal fibrosis was increased in right PPS joints (1.33 ± 0.47) (Figure 6A) and right control joints (2.33 ± 0.47) (Figure 6D). Right PPS and control joints were statistically different (P=0.014).

Light microscopy of articular cartilage: Histologic evaluation of sample C1 and C3 via H&E revealed a significant difference in OA score for the right control joints compared to its left normal control joints (Figures 7A-7F) (P<0.002). C1 histologic score revealed the same damage as C3. In the right control joints, the lesion (C1 and C3:5.67 ± 0.47), was characterized by denudation, complete erosion of hyaline cartilage to level of subchondral bone accompanied by granulation tissue (Figure 7E). Some samples had changes in the contour of the articular surface characterized by microfractures (Figure 7F), deep layer cyst formation (Figure 7B) and reparative granulation tissue (Figure 7D) or fibrocartilage (Figure 7A) focally above the level of the eroded and denuded articular surface. In right PPS joints (C1 1.5 ± 0.50 and C3:2.0 ± 0.0), defects had a well-developed layer of typical articular cartilage containing chondrocytes in the deep and middle zone however...
the superficial zone contained more fibrous layer of the flattened fibrocartilaginous cells. Chondrocytes deep within the cartilage layer had continued to undergo the typical sequence of differentiation (Figures 7G-7J). There was evidence of replacement of the dead chondrocytes and resorption and reconstitution of the extracellular matrix within the repair tissue (Figures 7K and 7L). Sections with partial defects (vertical fissures) had no healing evidence in control group (Figure 7C) however in PPS group; there was an evidence of minimal extra cellular hyaline-like matrix developed between fissures (Figure 7H). The right joint was significantly different in histological score between PPS and control groups (P=0.003).

**Scanning electron microscopy of the synovial membrane:** The synovial membrane of right joints revealed changes in both PPS (2 ± 0) and control group (4 ± 0). However control group was characterized by loss of synovial villi, deposition of cartilage fragments and formation of cuboidal crystals adhering to the synovial membrane (Figure 8A). Right PPS joints shows mild changes includes fibrosis but with presence of synovial villi in some regions (Figure 8B). There were significant differences in synovial villi destruction between the both groups (P=0.001).

**Scanning electron microscopy of the cartilage surface:** C2 in
right control joints had partial and full thickness cartilage fissures passed through the articular surface and penetrated deeply into the radial zone and sometimes to the level of the subchondral bone (4 ± 0) (Figures 9A-9C). However in right PPS treated joints these defects were fully filled with repair tissue (1 ± 0) (Figure 9D). This repair tissue was rough and acellular however in some areas, evidence of chondrocytes is revealed by the occurrence of pits. There was a significant difference between PPS and control right joints (P=0.001). C4 in right control joints revealed the same damage as C2 (Figures 9B and 9C). However in right PPS-treated joints (2 ± 0) large (Figure 9E) and small fissures (Figure 9F) were partially filled with repair tissue and the remaining lesions were limited to partial thickness fissures in some locations.

Discussion

In the present study, injection of allogenous cartilage particles (ACP) effectively resulted in clinical, gross, and histologic changes indicative of osteoarthritis. The lesions characterized by both chronic synovitis and osteoarthritis that includes partial and full-thickness cartilage erosion. During this study, no adverse effects were recorded and a mild degree of lameness was induced.

Clinically, NaPPS has been used to treat horses with mild or early-stage osteoarthritis, particularly with multiple joint involvements, because it is a systemic treatment rather than an intra-articular one. The ACP model used in the present study creates marked osteoarthritis and should be ideal to evaluate the efficacy of NaPPS.

In the present study, the dosage administered (3 mg/kg, once weekly for 4 weeks) was effective and elicit a therapeutic response without adverse effects in all clinical variables in this model. This is in contrast to another study [19] in which the author mentioned that...
the same dose was too low to elicit a therapeutic response but this difference may be due to the nature of the experimental animal model.

Improvement in the synovial membrane histological score and reduction of cartilage fragments size in PPS treated joints indicates that PPS enhances the phagocytic activities of the mononuclear leukocytes to engulf the foreign bodies. Furthermore, reduction in synovial membrane vascularity confined to the anti-angiogenesis effect of PPS.

Deposition of CPPD or BCP synovial crystals in osteoarthritis are common in human [31] and equines [32] clinical cases. This crystal is a consequence of rising of Ca, P and Mg [33-35] ions however, the mechanism of its formation is still under investigation.

Rising of the Ca, Mg and P ions are consequently a reason of broken tidemark and exposure of the subchondral bone [24].

Formation of large russet shape crystals was a feature of the ACP.
model [24] however in the present study formation of small cuboidal crystals in the control group. This difference may be due to decrease of the Ca concentration on the synovial fluid after cessation of ACP injection.

The cuboidal crystals are formed due to rising of P and Mg concentration on the control group however it is not found on the PPS group due to lowering of its concentration with PPS treatment.

PPS decrease crystals formation through repairing of the broken tide mark and enhancing the healing of the cartilage upper layer. This leads to dissolution of the crystals and diffuse out of the Ca, P and mg.

Using Pritzker grading system was suitable for evaluating the osteoarthritic joints however; in the evaluation of the healing stage, it may need some addition of healing features to grade 1, 2, 3 and 4.

This study confirms the previous reports on repair of articular cartilage in the horses. Among these studies, it is generally accepted that articular cartilage has limited reparative powers [36,37].

Neochondrogenesis can be stimulated by PPS as healing of full-thickness cartilage defects benefits from the production of granulation issue by subchondral mesenchymal cells. This tissue matures with time and undergoes metaplasia to fibrocartilage, or hyaline-like cartilage [36].

Convalescence time has been the subject of considerable debate. Granulation tissue was reported to fill the defects in one month [38], however maturation to fibrocartilage varies from 6 weeks [39], 3 month [40] and four month [38,41,42]. Development of the hyaline like cartilage is also varies from 8 weeks [39], 4 month [43], 6 month [38], 12 month [41,44] and 15 month [42].

This study reported that at 56-day post treatment, control joints with full thickness cartilage defects were filled with fibrocartilage. However the PPS treated joints were filled with hyaline-like tissue and chondroblasts attached to the subchondral bone and fibrous-like tissue superficially. Complete healing was not achieved in any treatment group.

Variation on healing stage in individual animals or regions in the same region influenced by influenced by a number of factors including the thickness, size, and anatomical location of the defect [36]. Post-traumatic and postoperative management of joint injuries varies considerably and is poorly understood. However, the degree of weight-bearing allowed and the length of convalescence can affect the quality of the reparative tissue [36,37].

In the present study, defects of healing confined to some zones may be due to presence of the old calcified cartilage layer, which may interfere with the attachment of the repair tissue to the subchondral bone [44].

This study confirm the previous morphological studies documented the inability of superficial or partial-thickness cartilaginous defects to undergo repair. Since the inflammatory response is absent, repair is limited to intrinsic healing which has limited potential for complete repair [36].

Previous report studying healing of superficial cartilage defects with SEM reported no healing 6-month post defect creation [45]. However, long term study (18 months) reported homogeneous material with a pitted-like appearance on the floor of the defects somewhat similar to that of surrounding cartilage but the surface of the defect was invariably quite rough and irregular [46]. Our study is the first to examine the healing of partial and full thickness cartilage defects. Examination of the articular cartilage with SEM was able to detect improvement in PPS joints. Disappearance of cartilage erosion in PPS treated joints and the appearance of a pitted surface suggests that a newly layer of matrix is largely formed. It is interesting to note that as long as an acellular layer of matrix persists no pits are seen but when such a layer presumably disappears or becomes attenuated the presence of the underlying chondrocytes is revealed by the occurrence of pits. This is in keeping with the idea that pits as seen on the surface of normal articular cartilage are due to the presence of superficially placed chondrocytes.

PPS modulates proteolytic enzymes in vitro and protects cartilage from proteoglycan loss from the extracellular matrix. This anticoncatabolic effect, combined with the antithrombotic effect, may result in improved cellular nutrition and matrix homeostasis [7]. These features would be supported by the significant decrease in articular cartilage fibrillation with NAPPS treatment observed in the present study.

These data clearly demonstrate the ability of PPS to stimulate the biosynthesis of components of the extracellular matrix accompanied by limiting their degradation by its anticoncatabolic effects. These beneficial pharmacological activities of PPS have resulted in its widespread use for the treatment of osteoarthritis in both veterinary [8,10,11,12], and human practice [14-16]. Moreover, the present study confirm previous works demonstrating the ability of this drug to induce chondrogenic differentiation of mesenchymal progenitor stem cell in vivo [47] and in vitro [48].

Authors’ Declaration of Interests

There are no conflicts of interest.

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Author Contributions Statement

All authors have made substantial contributions to all three of sections below:

- The conception and design of the study, acquisition of data, analysis and interpretation of data.
- Drafting the article or revising it critically for important intellectual content
- Final approval of the version to be submitted.

Ahmed M Elmesiry is the author responsible for the integrity of the work as a whole, from inception to finished article.

Ethical Board Review Statement

The experiment was approved by the Committee on Animal Experimentation at the Kafrelsheikh University, Egypt (Protocol no. VET1789 - 1/4/2013).

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