

Therapeutic Efficacy and Safety of Undenatured Type II Collagen Singly or in Combination with Glucosamine and Chondroitin in Arthritic Dogs

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ABSTRACT This investigation was undertaken to evaluate the therapeutic efficacy and safety of glycosylated undenatured type II collagen (UC-II) alone or in combination with glucosamine HCl and chondroitin sulfate in arthritic dogs. Twenty dogs divided into four groups ($n = 5$) were daily treated orally for 120 days: group I, placebo; group II, 10 mg UC-II; group III, 2,000 mg glucosamine + 1,600 mg chondroitin; group IV, UC-II (10 mg) + glucosamine (2,000 mg) + chondroitin (1,600 mg), followed by a 30-day withdrawal period. On a monthly basis, dogs were examined for overall pain, pain upon limb manipulation, and exercise-associated lameness. Serum samples were analyzed for markers of liver function (ALT and bilirubin) and renal function (BUN and creatinine). Body weight was also measured at a monthly interval. Dogs in group I exhibited no change in arthritic conditions. Dogs receiving UC-II alone showed significant reductions in overall pain within 30 days (33%) and pain upon limb manipulation and exercise-associated lameness after 60 days (66% and 44%, respectively) of treatment. Maximum reductions in pain were noted after 120 days of treatment (overall pain reduction, 62%; pain reduction upon limb manipulation, 91%; and reduction in exercise-associated lameness, 78%). The overall activity of the dogs in the UC-II supplemented with glucosamine and chondroitin group (group IV) was significantly better than the glucosamine + chondroitin-supplemented group (group III). Glucosamine and chondroitin alleviated some pain, but in combination with UC-II (group IV) provided significant reductions in overall pain (57%), pain upon limb manipulation (53%), and exercise-associated lameness (53%). Following withdrawal of supplements, all dogs (groups II to IV) experienced a relapse of pain. None of the dogs in any groups showed any adverse effects or change in liver or kidney function markers or body weight. Data of this placebo-controlled study demonstrate that daily treatment of arthritic dogs with UC-II alone or in combination with glucosamine and chondroitin markedly alleviates arthritic-associated pain, and these supplements are well tolerated as no side effects were noted.

KEYWORDS UC-II; Glucosamine; Chondroitin; Dog Arthritis; Pain Measurement

INTRODUCTION

Arthritis is a chronic disease that commonly affects large-breed dogs due to overweight/obesity, lack of exercise, physical injury, infection of joint surfaces, immune disorder, aging, or genetic predisposition. In particular, overweight and obesity in dogs can indirectly influence the degenerative joint disease process by increasing joint stress (Richardson et al. 1997). Dogs suffer most often with osteoarthritis than with rheumatoid arthritis (Hielm-Bjorkman et al. 2003). Arthritis is one of the most prevalent chronic health problems in the United States, affecting nearly 43 million people (Helmick et al. 1995). Although arthritis is often thought of as a disease that predominantly affects the elderly, it is the number one cause of disability affecting those over the age of 15. In fact, more than half of those affected by arthritis are under the age of 65, and almost 300,000 of those affected are children (Centers for Control and Prevention [CDC] 1994; Helmick et al. 1995; Trentham et al. 2001). Each year, arthritis is responsible for 44 million outpatient visits and almost 1 million hospitalizations, and is second only to heart disease in terms of its effect on disability from work (CDC 1994). As might be imagined from these statistics, the toll that arthritis takes on the health care industry is substantial, costing the United States approximately 65 billion dollars each year in health-related expenses. Unfortunately, the prevalence of arthritis does not appear to be decreasing, and by the year 2020 the CDC predicts that almost 60 million Americans will suffer from some form of arthritis.

Osteoarthritis is an inflammatory joint disease characterized by degeneration of the cartilage, hypertrophy of bone at the margins, and changes in the synovial membrane, and that eventually results in pain and stiffness of joints (Goldring 2000; Peat et al. 2001; Bellamy et al. 2001). Rheumatoid arthritis is a chronic disease characterized by inflammation, pain, swelling, and stiffness of multiple joints (Trentham et al. 1993; Okada 2000; Aceves-Avila et al. 2001). In either form of arthritis, dogs usually limp and are unable to move normally. Dog owners and veterinarians rarely notice the early warning signs of arthritis in dogs, since the dogs have the tendency to ignore soreness and discomfort until the arthritic signs are progressed significantly. Together, osteoarthritis and obesity cause a decreased quality of life for pets since joint pain is strongly associated with body weight (Richardson et al. 1997).

The present therapy of arthritis in dogs relies upon drugs that alleviate pain, control inflammation, and preserve ability to perform daily activity. Chronic use of cyclooxygenase (COX) inhibitors (nonsteroidal anti-inflammatory drugs [NSAIDs]) is linked to numerous side effects, including gastrointestinal (GI) bleeding and hepatic and renal dysfunction (Physician's Desk Reference [PDR] 1998; Muhlfeld and Floege 2005). Anti-inflammatory drugs, such as aspirin and ibuprofen, are nonspecific inhibitors of COX enzymes (both COX-I and COX-II), and they inhibit the production of inflammatory prostaglandins, thereby providing therapeutic effect, but they also inhibit the production of constitutive prostaglandins, causing severe side effects, such as severe GI bleeding (Matteson 2000). In the recent past, two commonly used drugs approved by the Food and Drug Administration (FDA) in arthritic dogs included Rimadyl (carprofen) and Deramaxx (deracoxib). Both Rimadyl and Deramaxx are NSAIDs. Rimadyl is not recommended for animals with known bleeding disorders and should not be used if a dog has pre-existing liver disease, inflammatory bowel disease, or a known tendency toward GI ulceration. Labrador Retrievers and other breeds are represented in the population that has experienced side effects or a fatal outcome from Rimadyl. Deramaxx (deracoxib), a COX-II inhibitor similar to Celebrex (celecoxib) and Vioxx (rofecoxib), prescribed as pain relievers for people, was withdrawn from the market in 2004, because of heart attack and stroke risk. According to a recent clinical study from the Mayo Clinic, other rheumatoid arthritic drugs such as Humira (adalimumab) or Remicade (infliximab) may cause serious infections or lead to the development of several kinds of cancer (Bongartz et al. 2006). Therefore, under the present circumstances, a safe therapy is needed for arthritic dogs.

In recent years, due to widespread availability of nutraceuticals, glucosamine and chondroitin sulfate were the two most commonly used supplements to ease the pain and discomfort of arthritis in dogs. Nutraceuticals are defined as functional foods, natural products, or parts of food that provide medicinal, therapeutic, or health benefits, including the prevention or treatment of disease. Glucosamine is an amino-monosaccharide precursor of the disaccharide unit of glycosaminoglycan, which is the building block of proteoglycans, the ground substance of articular cartilage (Paroli et al. 1991). Chondroitin sulfate is a part of a large protein molecule (proteoglycan) that

gives cartilage elasticity. Glucosamine is extracted from crab, lobster, or shrimp shells, and chondroitin is extracted from animal cartilage, such as tracheas and shark cartilage. In a recent pilot study, we found for the first time that daily administration of glycosylated undenatured type II collagen (40 mg of UC-II providing 10 mg/day) for 90 days significantly ameliorated the signs and symptoms of arthritis in dogs (DeParle et al. 2005). UC-II is a glycoprotein from chicken sternum cartilage. The presence of glycosylated “active” epitopes in the UC-II collagen matrix was confirmed by a validated ELISA method available from ChronoDEX, LLC (ArthroGen-CIA Capture “ELISA” test). Furthermore, electron microscopic analysis of UC-II was conducted to demonstrate the conformational integrity of the undenatured triple helical structure of a protein (Bagchi et al. 2002). Based on this study and other research (Bagchi et al. 2002), the present investigation was carried out to evaluate the therapeutic efficacy and safety of the antiarthritic compound UC-II (10 mg/day) alone or in a combination with the two most commonly used nutraceutical supplements (glucosamine HCl and chondroitin sulfate) in arthritic dogs given daily for 120 days, followed by a 30-day withdrawal. Glucosamine and chondroitin sulfate have become popular supplements for arthritis and are widely used to ease the pain and discomfort in arthritic dogs. Another objective of this investigation was to determine if the UC-II in combination with glucosamine and chondroitin was well tolerated by arthritic dogs.

MATERIALS AND METHODS

Animals

A group of 20 adult client-owned dogs was selected for this study based on the signs of arthritis, such as joint stiffness, lameness, and pain at the level of moderate severity. These dogs had swollen joints, and were experiencing difficulty in getting up or down (from a sitting and standing position) and walking (horizontal areas and short stairways). Arthritic dogs having any other serious disease or complications (such as hepatic or renal disease) were not included in the study. The owner consent was obtained before initiation of any experiments. The protocol of the present investigation for using arthritic dogs and their treatment was in compliance with the Murray State University Guidelines.

Supplements

UC-II is a standardized, undenatured (native) type II collagen complex containing 10 mg undenatured type II collagen in capsule form for use as a dietary supplement and was provided by InterHealth Nutraceuticals Inc., Benicia, CA, USA. UC-II is manufactured in a GMP facility at a low temperature, which preserves its undenatured form and biological activity. Glucosamine HCl and chondroitin sulfate were also provided as capsules by InterHealth Nutraceuticals.

Experimental

Twenty arthritic dogs were randomly divided into four groups (n=5) and received daily treatment as follows: group I (placebo), group II (10 mg UC-II), group III (2,000 mg glucosamine HCl + 1,600 mg chondroitin sulfate), and group IV (10 mg UC-II + 2,000 mg glucosamine HCl + 1,600 mg chondroitin sulfate). The treatment was given daily for 120 days, followed by a 30-day withdrawal period. None of the dogs received any NSAIDs for 3 to 4 weeks before the study or during the study period. The study was conducted double-blinded; that is, the investigators or owners had no knowledge of the capsule contents.

Pain and Body Weight Measurements

The dogs were evaluated for overall pain, pain upon limb manipulation, and exercise-associated lameness on a monthly basis for a period of 150 days. Overall pain was measured as a general gross observation, which included trouble in standing after sitting, or trouble in sitting after standing, vocalization, crying, etc. Results were graded on a scale of 0 to 10: 0, no pain; 5, moderate pain; and 10, severe and constant pain. Pain upon limb manipulation was evaluated by animals' vocalization or other observations of pain during the extension and flexion of all four limbs for few minutes. Results were graded on a scale of 0 to 4: 0, no pain; 1, mild; 2, moderate; 3, severe; and 4, severe and constant. Lameness was measured after physical exercise for gross observations, which included limping, holding limb up, rigidity of limbs, etc. Signs of pain and lameness were noted on a scale of 0 to 4: 0, no pain; 1, mild; 2, moderate; 3, severe; 4, severe and constant. Severity of pain during various activities, such as standing from sitting, sitting from standing, playing, and vocalization, and during extension and

flexion of limbs was the basis for gradation. Body weights and physical evaluation were determined on a monthly basis. Gross observations were evaluated and recorded monthly using a questionnaire regarding the overall activity and the improvement of the arthritic symptoms. Furthermore, overall performance of individual groups was assessed, which include running, participation in jogging activities, movement up and down stairs, comfort ability in moving from sitting to standing position(s), cheerful attitude toward playing and jumping, etc.

Biochemical Assays

Blood samples were collected by jugular venipuncture using 22-gauge needles and 12-mL syringes. Serum was separated in a marble-top tube (serum separating tubes without anticoagulant) and transferred into plastic snap-top tubes. Serum samples were frozen immediately and kept at -80°C until analyzed for blood urea nitrogen (BUN), creatinine, bilirubin, and alanine aminotransferase (ALT), using Beckman Coulter CX5-PRO Synchron Clinical System (Fullerton, CA). Bilirubin and ALT were used as markers of liver function and BUN and creatinine were used as markers of renal and heart function.

Statistical Analysis

The data of serum chemistry in Table 2 and of pain observation in Figures 1 to 3 are presented as mean $\pm$ SEM. Statistical significance of differences was determined by analysis of variance (ANOVA) coupled with Tukey-Kramer test using the NCSS 2000 Statistical System for Windows. Differences with $p < 0.05$ were considered statistically significant.

RESULTS

Data of pain evaluation in arthritic dogs receiving placebo and those receiving UC-II alone or in combination with glucosamine and chondroitin are shown in Figures 1 to 3. Dogs receiving placebo (group I) exhibited no significant change in arthritic conditions at any time during the course of treatment. Dogs receiving UC-II (10 mg/day) alone (group II) showed significant reduction in overall pain within 30 days (33%) and pain upon limb manipulation and exercise-associated lameness after 60 days (66% and 44%, respectively) of treatment. Maximum reductions

in the pain were noted after 120 days treatment (overall pain, 62%; pain upon limb manipulation, 91%; and exercise-associated lameness, 78%). Glucosamine plus chondroitin (group III) alleviated some pain but not significantly ($p > 0.05$). Group IV dogs receiving a combination of UC-II and glucosamine plus chondroitin showed marked reductions in overall pain (57%), pain upon limb manipulation (53%), and exercise-associated lameness (53%) after a daily treatment of 120 days. Following the withdrawal of supplements in either of the groups (group II to IV) for 30 days, all dogs experienced a relapse of overall pain, pain upon limb manipulation, and exercise-related lameness. In addition, UC-II-treated dogs were more playful, energetic, and less painful compared to other groups used in this study.

None of the dogs receiving dietary supplements showed any signs of adverse effects. There were no significant changes in any markers of liver function (ALT and bilirubin) or renal or heart function (BUN and creatinine) during the course of this investigation. Body weight remained within the normal range throughout the course of this study. An increase in body weight was observed in the placebo group, demonstrating that activity level in that group was less than the other supplemented group (Table 1).

DISCUSSION

This investigation was pursued with two specific objectives: (a) whether the combination of UC-II and glucosamine plus chondroitin provides better antiarthritic effects than glucosamine plus chondroitin or UC-II alone, and (b) whether these supplements are well tolerated by arthritic dogs, following a long term of their use. The present findings revealed that UC-II therapy (10 mg/day) alone or in combination with glucosamine plus chondroitin for 120 days provided significant improvement in the overall pain, pain upon limb manipulation, and pain after physical exertion. The greatest physical improvements were noted in UC II supplemented group after a treatment period of 120 days, suggesting that prolonged treatment with the supplement may lead to better therapeutic results. After a 30-day withdrawal period, all dogs who had received treatment suffered from a relapse of signs and symptoms associated with arthritic conditions, such as pain and lameness. All supplements were well tolerated and no adverse effects were observed.

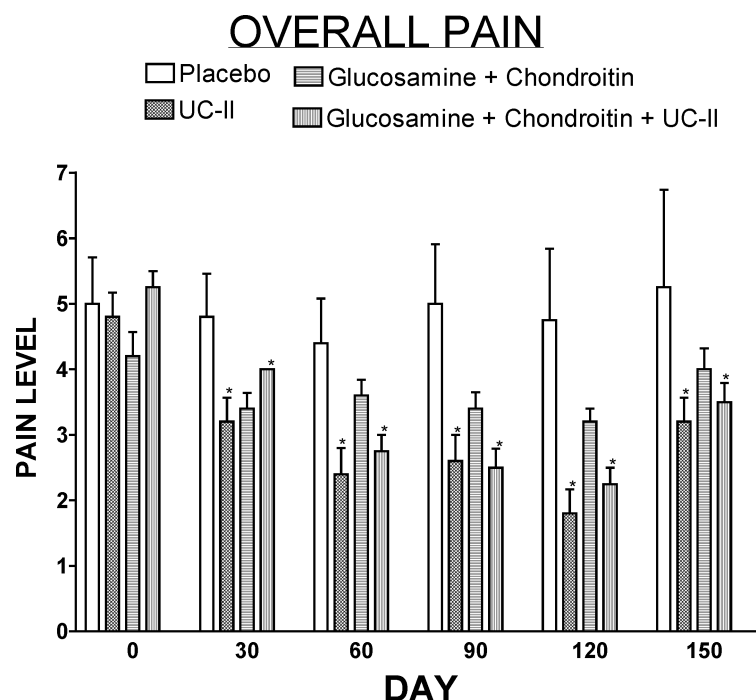


FIGURE 1 Effects of UC-II (10 mg) and/or glucosamine HCl (2,000 mg) plus chondroitin sulfate (1,600 mg) given daily for 120 days, followed by a 30-day withdrawal period, on overall pain in arthritic dogs. Overall pain was graded on a scale of 0 to 10: 0, no pain; 5, moderate pain; and 10, severe and constant pain. * Indicates significant difference from pretreated values ($p < 0.05$).

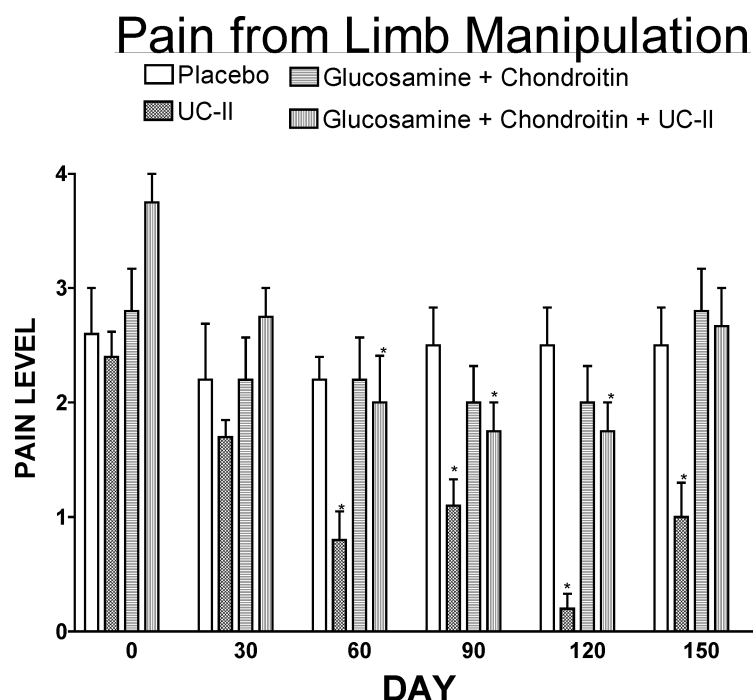


FIGURE 2 Effects of UC-II (10 mg) and/or glucosamine HCl (2,000 mg) plus chondroitin sulfate (1,600 mg) given daily for 120 days, followed by a 30-day withdrawal period, on pain upon limb manipulation in arthritic dogs. Pain upon limb manipulation was evaluated by animal's vocalization or other observations of pain during the extension and flexion of all four limbs for few min. Results were graded on a scale of 0 to 4: 0, no pain; 1, mild; 2, moderate; 3, severe; and 4, severe and constant. * Indicates significant difference from pretreated values ($p < 0.05$).

TABLE 1 Effects of UC-II (10 mg) and/or glucosamine HCl (2,000 mg) plus chondroitin sulfate (1,600 mg) given daily for 120 days, followed by a 30-day withdrawal period, on body weight (pounds) of dogs

Day	Placebo	UC-II	Glucosamine + Chondroitin	Glucosamine + Chondroitin + UC-II
0	69.76 ± 11.49 (100)	62.25 ± 6.29(100)	73.20 ± 7.47 (100)	59.28 ± 9.66 (100)
30	68.88 ± 10.95 (99)	62.35 ± 6.38 (100)	72.80 ± 7.88 (100)	58.83 ± 8.90 (99)
60	69.84 ± 10.86 (100)	62.33 ± 6.55 (100)	73.12 ± 7.31 (100)	57.53 ± 10.04 (96)
90	76.80 ± 10.20 (110)	62.32 ± 6.97 (100)	72.96 ± 7.39 (100)	58.70 ± 9.18 (99)
120	79.58 ± 12.65 (114)	62.83 ± 6.71 (101)	76.16 ± 8.04 (104)	59.53 ± 8.74 (100)

Values are means ± SEM (n = 4–5).

Note: Body weight values remained significantly indifferent throughout the study period compared to pretreated values ($p > 0.05$).

In a recent double-blind pilot study, UC-II (1 mg or 10 mg/day, PO for 90 days) was found to be significantly effective in ameliorating arthritic pain in dogs (DeParle et al. 2005). A 10-mg dose of UC-II provided markedly greater effects in all parameters measured for the arthritic symptoms than 1 mg dose in improving the overall performance and well-being of the dog. The findings of the present study revealed that the arthritic dogs receiving UC-II (10 mg/day) in combination with glucosamine and chondroitin (group IV) overall performed better than glucosamine (2,000 mg) plus chondroitin (1,600 mg) (group III). The

most commonly used two nutraceuticals (glucosamine and chondroitin) provided some beneficial antiarthritic effects, but not significantly ($p > 0.05$), which suggested that the observed therapeutic effects were mainly due to the supplementation of UC-II. The findings also revealed that though the beneficial effects of UC-II were observed within 30 days, maximum effects were seen after 120 days. UC-II functions through a process called oral tolerization (i.e., this process takes place in the small intestine, where food is absorbed). Through a complex series of immunological events, patches of lymphoid tissue (Peyer's patches) surrounding the small

TABLE 2 Effects of UC-II (10 mg) and/or glucosamine HCl (2,000 mg) plus chondroitin sulfate (1,600 mg) given daily for 120 days, followed by a 30-day withdrawal period, on markers of liver and renal functions in serum of dogs

	DAY					
	0	30	60	90	120	150
ALT						
Group I	26.00 ± 3.33	22.20 ± 1.98	22.00 ± 1.97	23.75 ± 2.20	22.50 ± 1.66	23.50 ± 1.55
Group II	29.00 ± 2.57	29.00 ± 2.41	30.20 ± 4.08	28.80 ± 3.99	25.20 ± 2.78	25.00 ± 2.19
Group III	21.75 ± 6.97	22.20 ± 3.97	21.00 ± 4.05	19.60 ± 2.75	21.60 ± 2.80	27.20 ± 5.60
Group IV	22.80 ± 3.89	21.50 ± 2.20	23.78 ± 4.82	20.00 ± 4.99	18.25 ± 3.29	23.00 ± 4.82
BILIRUBIN						
Group I	0.52 ± 0.12	0.60 ± 0.16	0.68 ± 0.08	0.43 ± 0.18	0.56 ± 0.09	0.52 ± 0.13
Group II	0.46 ± 0.11	0.52 ± 0.17	0.50 ± 0.08	0.60 ± 0.13	0.70 ± 0.17	0.42 ± 0.13
Group III	0.46 ± 0.12	0.62 ± 0.21	0.40 ± 0.07	0.62 ± 0.14	0.42 ± 0.07	0.50 ± 0.15
Group IV	0.56 ± 0.14	0.48 ± 0.06	0.48 ± 0.03	0.75 ± 0.19	0.50 ± 0.11	0.49 ± 0.06
BUN						
Group I	16.80 ± 1.24	18.00 ± 2.07	13.20 ± 1.65	21.50 ± 1.66	19.00 ± 3.44	17.50 ± 1.32
Group II	17.80 ± 2.65	18.80 ± 2.85	18.40 ± 0.51	18.40 ± 4.72	18.80 ± 1.96	14.40 ± 1.43
Group III	15.20 ± 1.83	16.80 ± 2.10	19.20 ± 2.40	19.60 ± 3.91	15.60 ± 2.04	17.00 ± 1.87
Group IV	19.40 ± 2.58	24.00 ± 2.81	15.75 ± 1.70	21.25 ± 2.29	18.50 ± 3.52	16.00 ± 2.64
CREATININE						
Group I	0.94 ± 0.06	0.92 ± 0.05	0.90 ± 0.03	0.88 ± 0.05	0.87 ± 0.05	0.92 ± 0.05
Group II	1.06 ± 0.07	0.96 ± 0.08	0.98 ± 0.04	0.90 ± 0.08	0.94 ± 0.10	0.96 ± 0.07
Group III	0.92 ± 0.07	0.96 ± 0.05	1.00 ± 0.05	0.94 ± 0.10	0.92 ± 0.07	0.90 ± 0.04
Group IV	0.86 ± 0.02	1.12 ± 0.25	0.80 ± 0.07	0.95 ± 0.07	0.90 ± 0.07	0.87 ± 0.06

Values are means ± SEM (n = 5).

Note: Values of markers of liver function (ALT and bilirubin) and renal function (BUN and creatinine) remained significantly indifferent throughout the study period compared to pretreated values ($p > 0.05$).

Pain After Physical Exertion

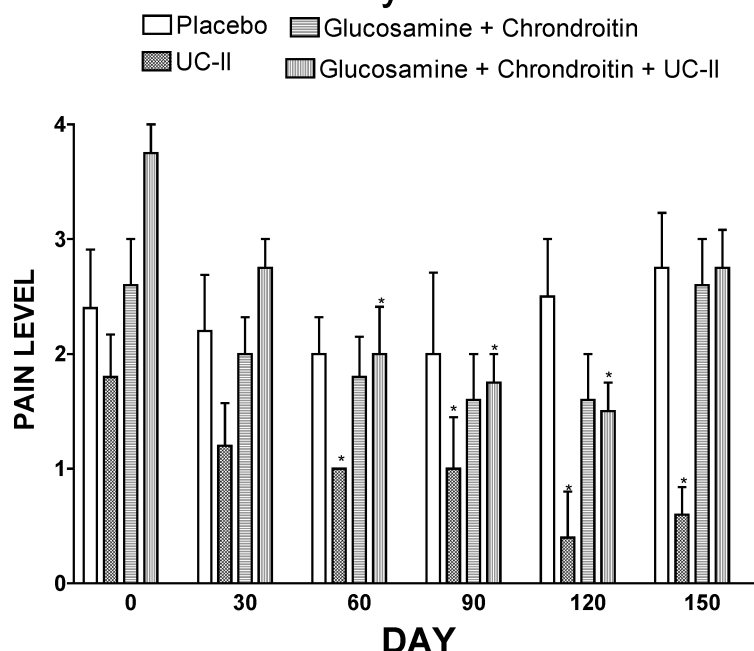


FIGURE 3 Effects of UC-II (10 mg) and/or glucosamine HCl (2,000 mg) plus chondroitin sulfate (1,600 mg) given daily for 120 days, followed by a 30-day withdrawal period, on pain after physical exercise in arthritic dogs. Lameness was measured after physical exercise for limping, holding limb up, rigidity of limbs, etc. Signs of pain and lameness were graded on the scale of 0 to 4: 0, no pain; 1, mild; 2, moderate; 3, severe; and 4, severe and constant. * Indicates significant difference from pretreated values ($p < 0.05$).

intestine screen incoming compounds and serve as a “switch” to turn the body’s immune response to foreign substances on or off, depending on what the substance is (Mowat 1987). In the case of type II collagen, small amounts of UC-II (typically 10 mg or less/day, PO) have been shown to turn off the immune response targeted at type II collagen present in bone joint cartilage without any side effects (Weiner et al. 1994; Sieper et al. 1996; Gimsa et al. 1997). This process helps the body to differentiate between elements that are foreign invaders to the body and those that are nutrients (Weiner 1997; Trentham 1998). Previous studies have demonstrated that UC-II improves joint mobility and flexibility by the mechanism of preventing the immune system from attacking and damaging its own joint cartilage (Mowat 1987; Sieper et al. 1996; Gimsa et al. 1997; Trentham 1998; Trentham et al. 1993, 2001).

Unlike UC-II, glucosamine and chondroitin are expected to cause decrease of pain sensation, provide an improved resistance to additional joint tissue breakdown, and rejuvenate some joint tissues. Studies suggest that glucosamine helps to relieve pain by enhancing proteoglycan synthesis, which is impaired in osteoarthritic cartilage (Hougee et al. 2006). Chon-

droitin sulfate aids in keeping cartilage tissue from dehydrating and assists in cushioning impact stress and reduce joint pain. Earlier studies demonstrated that combining chondroitin sulfate with glucosamine may improve arthritic symptoms and may have some beneficial effects. However, these supplements are not known to reverse structural changes in a joint such as torn cartilage, calcium deposits, and advanced scar tissue. In spite of these beneficial effects of glucosamine and chondroitin described elsewhere, in the present study we did not find significant antiarthritic effects in dogs using the above combination. A recent multicenter with 1,583 patients with symptomatic osteoarthritis in the knee received 1,500 mg of glucosamine and 1,200 mg of chondroitin sulfate daily. They reported that the combination or supplements alone did not reduce pain effectively in the overall treatment groups (Clegg et al. 2006).

Another benefit of the present study resulted in healthy weight maintenance in supplemented groups, compared to control. Body weight was maintained in all supplemented groups, while there was an increase in body weight in the placebo group. This demonstrated that the dogs were more active and arthritic symptoms were remarkably reduced in the supplemented group.

Weight control has been shown to indirectly influence the degenerative joint disease process by reducing the stress on the joint (Eaton 2004). A small amount of weight loss reduces the risk of developing arthritis in dogs and human.

In conclusion, the present study suggests that daily treatment of arthritic dogs with UC-II alone or in combination with glucosamine plus chondroitin ameliorates signs and symptoms of arthritis significantly greater than glucosamine and chondroitin. The results of this study also demonstrate that the supplements are well tolerated. The relapse seen in dogs 30 days posttreatment showed that continuous treatment is required.

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