4 courses, and 760 (1.5%) received 5 courses or more. The time from diagnosis to when 50% of the subset received TKR was significantly longer (p1 year, an average of 9 months longer than those with no IA HA. Patients who received \geq 5 courses had a delay in TKR by 3.6 years. **Conclusions:** Among 182,022 patients with knee OA, those who received IA HA had a significantly longer time before TKR. More courses of IA HA injections were associated with a longer time to TKR. This study suggests a significant clinical benefit from use of IA HA for OA as delay in time to TKR can have important clinical and economic implications.

Median Time from knee OA diagnosis to TKR by number of courses of IA HA

	No IA HA	1	2	3	4	5+
Days to TKR	114	386	648	875	1054	1312
Years to TKR	0.3	1.1	1.8	2.4	2.9	3.6

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EVALUATION OF THE CLINICAL EFFICACY OF AUTOLOGOUS CONDITIONED SERUM IN PATIENTS WITH COXARTHROSIS

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Purpose: The aim of this open study was a comparative assessment of the effectiveness of local therapy with autologous conditioned serum (ACS) and low molecular weight hyaluronic acid (LHA) in patients with coxarthrosis.

Methods: The study included 60 patients with firm coxarthrosis in accordance with ACR criteria at the age of 55,5 \pm 8,7. The main group (ACS treatment) consisted of 33 (55%) persons. Experimental group (LHA treatment) consisted of 27 patients who are comparable with the main group patients in terms of age, BMI, radiographic stage, disease duration and severity of clinical indicators. ACS was prepared in accordance with established method and injected intraarticularly (2.5 ml twice a week for three weeks). LHA treatment consisted of 3 weekly intraarticular injections of 40 mg sodium hyaluronate each. All intraarticular actions were performed with ultrasound control. Treatment efficiency was evaluated after 1.3 and 6 months after treatment, the following criteria were used: bodily pain dynamics in accordance with VAS, morning stiffness module and Womac index functional scale, and Lequesne index. "Area under the curve" (AUC) approach with estimation of treatment efficiency prolongation for 6 months (AUC6) was used for evaluation of clinical effect retention.

Results: A decrease in pain syndrome intensity according to VAS in hip joints at 1, 3 and 6 months of treatment in both compared groups was recorded.However, a significant regression was observed in the treatment of pain with ACS in comparison with LHA.After 1 month the decrease of pain severity in accordance with VAS was comparable (9.1%, p = 0.40), after 3 and 6 months, pain severity was higher in the LHA group compared with ACS group (+ 52.5%, p = 0.009 and + 33.1%, p = 0.047respectively). AUC6 in case of ACS treatment was 35,6% (p = 0,011) higher compared with LHA treatment. Extension of the clinical effectiveness expressed via "morning stiffness" module of the Womac index AUC6 indicator was 55.9% (p=0.003) higher in case of ACS treatment compared with LHA treatment. Womac functional scale in case of ACS improved through all three control timepoints (-23.1% -28.5% -39.1%; p = 0.001). Same dynamics was observed in LHA group as well (-35.6%, -26.4%; p = 0.001 and -20.4%; p = 0.005), however after 6 months of monitoring more significant improvement of articular function was recorded within ACS group compared with LHA (18.3%, p = 0.044). The overall clinical efficacy in accordance with Lequesne index with a six-month monitoring was comparable in both groups, the difference came up to 8,8% (p = 0,65). Conclusions: Intensity of pain, stiffness, functional status and overall clinical severity of coxarthrosis significantly and consistently decreased during therapy with ACS and LHA.However, favorable changes of local therapy with LHA were inferior in duration of conservation of treatment effect of ACS.

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REFINEMENT OF PRECLINICAL STUDIES FOR VISCOSUPPLEMENTATION THERAPY EVALUATION: INTEREST OF COMPLEMENTARY TECHNIQUES TO EVALUATE CARTILAGE IN A RABBIT MODEL OF EARLY OSTEOARTHRITIS

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Purpose: One of the major challenges of the viscosupplementation (VS) therapy is the development of more efficient formulations and, to this end, the use of animal models of osteoarthritis (OA) is still mandatory. The assessment of VS efficacy is challenging mainly since its structural effects are subtle. In order to refine preclinical VS efficacy studies and reduce the number of animals used, there is a crucial need for more sensitive and discriminant evaluation tools. In this study, we especially focused on complementary techniques to evaluate OA cartilage in a rabbit model of early OA.

Methods: Cranial cruciate ligament transection (ACLT) was performed in the left knee of white New-Zealand rabbits (n=12) to induce traumatic OA. One week post-ACLT and then weekly for 5 weeks, the operated knees of 6 rabbits were injected with a hyaluronic acid (HA) containing commercial formulation (Ostenil®, HA group). One group was injected with saline (operated-control group, n=6). The contralateral right knees (n=8) were used as unoperated-controls. Endpoint evaluation was done at the 6th week post-ACLT and included: gross and histological scoring of cartilage lesions, measurement of cartilage thickness by Equilibrium Partitioning Iodine Contrast micro-Computed Tomography (EPIC μ -CT) as well as the evaluation of the surface by Scanning Electron Microscopy (SEM).

Results: Gross and histological scorings showed statistical differences between operated and unoperated knees; however, no difference between the HA and operated-control groups was evidenced. SEM revealed that unoperated-control samples had normal smooth to rough surfaces with discreet cable-like structures. Cartilage from the operated knees presented rough surfaces and clearly visible cable-like structures, which might be the sign of cartilage matrix erosion. Finally, mean cartilage thickness and volume measured by EPIC µ-CT were comparable for the 3 groups. Interestingly, the use of the thickness distribution representation showed clear differences in the intact cartilage occurrences (i.e. thickness higher or equal to 0.9 mm). Indeed, intact cartilage proportion decreased statistically from 20% in the unoperated-control group to 10% in the operated-control group. In addition, intact cartilage proportion in the HA treated knees was equivalent to unoperatedcontrol group, highlighting a moderate efficacy of HA which was detected neither by histology nor by SEM.

Conclusions: Complementary techniques were implemented to evaluate cartilage lesions in a rabbit model of early OA in order to refine preclinical VS efficacy studies. This study points out that classical evaluation tools (macroscopy and histology) are sensitive enough to discriminate between cartilage from unoperated and operated groups in early OA but that EPIC- µCT also permits the detection of the subtle structural effects of HA, validating this technique as a powerful evaluation tool.

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EVIDENCE OF *IN VIVO* DRUG DELIVERY VIA THE TAT PROTEIN TRANSDUCTION DOMAIN

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Purpose: Drug delivery to synovial joints is a major limitation for developing effective treatments for osteoarthritis (OA). Synovial joint anatomy includes large pores that allow direct transport between synovial fluid and fenestrated capillaries. This transport preferentially selects larger molecules to remain within the joint and exports smaller