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**SPECIFIC DESTRUCTION OF THE JOINT STRUCTURE IN REACTIVE ARTHRITIS**

Axmedov Kh.S., Khalmetova F.I., Abdurakhimova L.A.

**ОСОБЕННОСТИ ДЕСТРУКЦИИ СУСТАВНОЙ СТРУКТУРЫ ПРИ РЕАКТИВНОМ АРТРИТЕ**

Ахмедов Х.С., Халметова Ф.И., Абдурахимова Л.А.

**REAKTIV ARTRITDA BO'G'IM STRUKTUR DESTRUKTSIYASINING O'ZIGA XOSLIGI**

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**Цель:** определение уровня белков матрикса хрящевых олигомеров у пациентов с реактивным артритом в зависимости от его клинической картины для определения ранней деструкции хряща. **Материал и методы:** в исследование включены 120 пациентов в возрасте  $36,9 \pm 12,3$  года, разделенных на три группы в зависимости от триггерных факторов, и 20 здоровых лиц. **Результаты:** на основании полученных данных о ранних стадиях реактивного артрита хламидийной этиологии, длительном течении и особенно повышении содержания хрящевого олигомерного матриксного белка у женщин можно сделать вывод об инициации ранней деструкции хряща в структуре сустава. **Выводы:** выявленные изменения указывают на активизацию изменений в структуре суставов и развитие вторичного остеоартроза на ранних стадиях заболевания.

**Ключевые слова:** реактивный артрит, олигомерный матриксный белок хряща, суставной синдром, деструкция, *Chlamydia trachomatis*.

**Maqsad:** reaktiv artritga chalingan bemorlarda tog'ayning erta destruksiyasini o'rganish. **Material va usullar:** tadqiqotga trigger omillarga ko'ra uch guruhga ajratilgan reaktiv artritga chalingan 120 nafar  $36,9 \pm 12,3$  yoshdagi bemorlar va nazorat guruhi sifatida 20 nafar sog'lom kishilar jalb qilingan. **Natijalar:** olingan natijalar asosida reaktiv artritning ilk davrlaridan xlamidiya etiologiyali, cho'ziluvchi kechishi va ayniqsa ayollarda tog'ay oligomer matriks oqsili ortib borishi bo'g'im strukturasiida tog'ayning erta destruksiyasi boshlanishi haqida xulosa qilish mumkin. **Xulosa:** bo'g'imlarda strukturali o'zgarishlar va kasallikning erta davrlaridan ikkilamchi osteoartrit shakllanishidan dalolat beradi.

**Kalit so'zlar:** reaktiv artrit, tog'ay oligomer matriks oqsili, bo'g'im sindromi, destruksiya, xlamidiya etiologiyasi.

Nowadays rheumatic diseases attract attention in all spheres of medicine. According to its prevalence reactive arthritis (ReA) occupies one of the leading places and according to various references it takes 4-40% [3,4]. Moreover, it is characterized by progression accompanied by articular syndrome and damage of major joints [10,13]. It leads to invalidation of young and middle-age patients, which has social significance and urgency [1,8,15].

It is known that, clinical presentation and articular syndrome of many rheumatologic diseases differ by specific morphologic alterations and inflammatory process [7,12]. Rheumatic diseases, undoubtedly, cause certain alterations due to different progression and variety of underlying mechanisms. According to various opinions, the factors vary from unfavorable environmental effects to profile of cytokines with negative effect on joints (TNF- $\alpha$ , IL-17, IL-23, IL-1, and IL-6). Moreover, ReA is considered to be a disease with different progression, which means it has different clinical and x-ray alterations, and functionally heterogeneous pathology [6,11,14]. According to the results of the performed study, articular lesions in ReA depend on its trigger factors. Especially urogenital ReA differs by active structural alterations and in 33.3% of the cases secondary osteoarthritis develops within the first year of the disease. In its turn, it causes activation of structural alterations, which leads to deterioration of patients' life quality. That is why, detection of early cartilage destruction in ReA has a practical importance.

Recently, according to the available data in literature, osseous erosions caused by proinflammatory cytokins in

ReA immune response (IL-1, IL-6 and TNF- $\alpha$ ) can serve the basis for degenerative alterations in cartilages. That process causes intensification of synthesis of collagenase and matrix metalloproteinase leading to splitting of 2 type of collagen. Results of the last scientific researches showed, that cartilage oligomere matrix protein (COMP) provides important information about metabolic changes occurring in cartilage matrix under the influence of the aforesaid enzymes. Consequently, rise of COMP in blood serum can serve as a biomarker of early cartilage destruction in ReA. That is why, study of change of COMP in blood serum dependently on the clinical progression of the disease in patients with ReA has scientific and practical value.

**The objective**

Study of dynamic changes of cartilage oligomere matrix protein (COMP) level in patients with ReA dependent on its clinical presentation for the definition of early cartilage destruction.

**Materials and research methods**

The study enrolled randomly selected 120 patients with average term of ReA equal to  $3.8 \pm 1.7$  years aged 18-50 ( $36.9 \pm 12.3$ ) years old. The major part of these patients were women, 70 (58.3%). For the study patients were classified to three groups according to etiological factors. The I group (n=60) had *Chlamydia trachomatis* etiology; the II group (n=30) had *Sinia enterocolitica*, and the III group (n=30) had *Campylobacter jejune* as an etiological factor. The control group involved 20 healthy subjects (average age  $37.5 \pm 6.2$  years old) with age and sex approximately compatible to patients with ReA (Table 1).

Clinical characteristics of patients with ReA, abs. (%)

Groups	Sex		Average age	Average term of the disease, years
	Male	Female		
1, n=60	25 (41.7)	35 (58.3)	32.9±11.1	2.9±1.8
2, n=30	13 (43.3)	17 (56.7)	36.4±6.8	3.1±1.9
3, n=30	12 (40)	18 (60)	34.1±7.3	2.8±1.9

For the detection of trigger infections we used immunologic (detection of antibodies to infectious antigen in blood serum), molecular-biological (polymerase chain reaction), and bacteriological tests. We had detection of infections causing ReA in all patients, after which they were examined by urologist, gynecologist (taking smear from urethra and vagina), and oculist.

For the assessment of the progression and activity of ReA we applied visual analogue scale (VAS), DAS and parameters of the acute stage of inflammation. Functional capabilities of the patients were assessed according to functional classes (FC), Health Assessment Questionnaire (HAQ) and Ritchie index. Laboratory tests included common blood analysis and biochemical blood tests. All patients had x-ray imaging of the joints. Cartilage oligomer matrix protein (COMP) and female sexual hormones were detected using enzyme immunoassay (ELISA, Russia).

Exclusion criteria were as follows:

- Patients with no confirmed diagnosis of ReA according to EULAR/ACR;
- Patients under 18;
- No surgical treatment of ReA within and before the period of study;
- Severe concomitant pathology (renal, hepatic, cardiac failure, high uncontrollable AH, decompensated diabetes mellitus, etc.); injures;
- Dangerous tumors, drinking of alcohol, psychic diseases, such as dementia and mental impairments;
- Patients with BMI below 29.

Statistical processing of the results was performed using Microsoft Office Excel 2013 software and standard statistical method.

### Results and discussion

Majority of the patients were 30-40 years old (69.1%) and women (58.3%). On the basis of the analysis of anamnesis morbid the average age of patients at the moment of appearance of initial symptoms of ReA

was 30.1±4.5 years old. Average time from the moment of appearance of initial symptoms till the diagnosis was 3.7 months.

The data in Table 1 show that, according to the form of the disease and etiological factors clinical presentation of ReA differs in three groups. There is reliable ( $p < 0.05$ ) difference of long-lasting progression of the pathology among the patients of the I group compared to other forms. At the same time, in 31.6% of the cases we could observe recurrent progression of urogenital ReA.

Dysfunction of joints can be linked with certain alterations relevant to inflammatory process. Indicators of acute stage, duration of morning stiffness and the number of inflamed joints in the patients of the I group caused limitation of functional capabilities of the joints. In their turn, patients of the II and III groups suffered monoarthritis, while the patients of the I group there were more cases of oligoarthritis and polyarthritis. It should be noted that, 100% of all patients in all three groups had synovitis, which was mostly singular and sometimes accompanied peri-arthritis manifested by tendinitis or bursitis.

It is known that, diseases proceeding with arthritis are characterized by certain alterations in joint structure. Immunologic alterations in ReA are based on synovial inflammation, disorders in its structure and development of fibrosis [5]. The aforesaid processes proceed with various alterations providing the possibility of bone erosion and articular surface incongruence. Perhaps, these transformations are linked with change in characteristics of cartilage morphologic substrate. So, rise of COMP in blood serum of the patients with ReA indicate metabolic changes in cartilage [9]. It should be noted that, changes of COMP level among the patients enrolled in the study varied greatly. As it is seen in Table 2, when compared to the control group, all male and female patients had a tendency for the rise of COMP, but the values were not statistically significant ( $p > 0.05$ ). However, in the I group the part of those who had rise of COMP reference values ( $> 1000$  ng/mL) was equal to 53.3% (Fig. 1).

Table 2

Clinical presentation of patients with ReA, abs. (%)

Symptoms	1 group, n=60	2 group, n=30	3 group, n=30
According to progression:			
- Acute	7 (11.7)	19 (63.3)	16 (53.3)
- Long-lasting	24 (40)	5 (16.7)	3 (10)
- Chronic	10 (16.7)	3 (10)	7 (23.3)
- Recurrent	19 (31.6)	3 (10)	4 (13.4)

Articular syndrome:			
- Monoarthritis	4 (6.6)	16 (53.3)	14 (46.7)
- Oligoarthritis	34 (56.7)	10 (33.4)	13 (43.3)
- Polyarthritis	22 (36.7)	4 (13.3)	3 (10)
- Sacroiliitis	53 (88.3)	26 (86.7)	4 (13.3)
- Spondylitis	40 (66.7)	4 (13.3)	13 (43.3)
- Dactylitis	6 (10)	5 (16.7)	6 (20)
Dysfunction of joints:			
- I class	12 (20)	18 (60)	15 (50)
- II class	29 (48.3)	8 (26.7)	11 (36.7)
- III class	19 (31.7)	4 (13.3)	4 (13.3)
Articular index and laboratory results:			
- Duration of morning stiffness, minutes	31.3±5.9	19.1±6.1	17.3±8.5
- Pain, VAS, mm	79.5±12.8	67.5±12.8	53.5±11.4
- Number of painful joints	8.2±3.7	4.1±0.7	4.1±0.7
- Number of swelling joints	5.6±0.6	2.5±2.6	2.1±0.9
- C-reactive protein, mg/L	19.8±3.9	12.8±1.9	11.8±1.7
- Erythrocyte sedimentation rate, mm/s	25.3±3.9	18.3±3.6	19.3±5.5

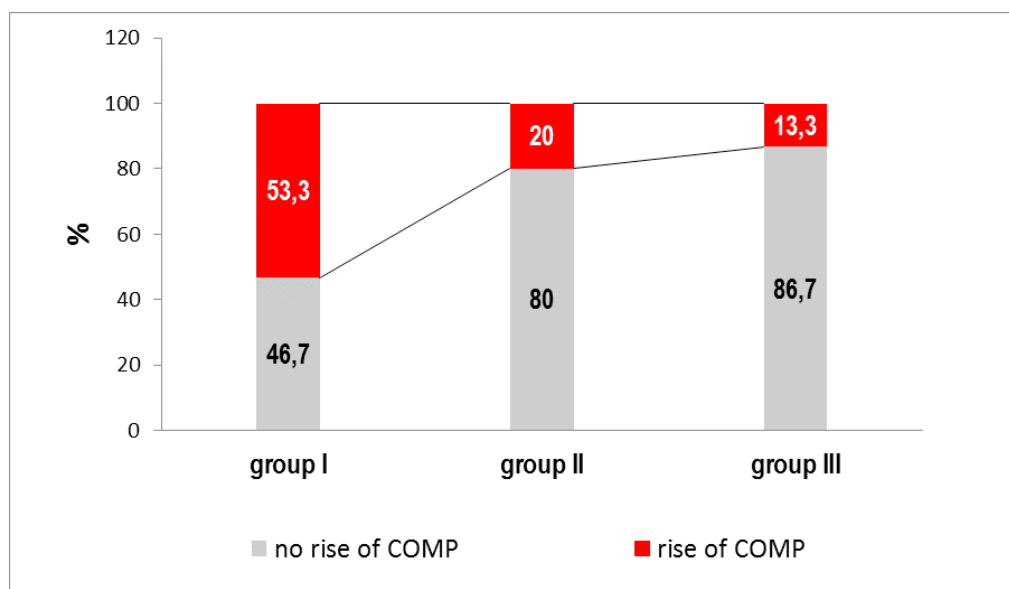


Fig. 1. COMP rise among the patients with ReA.

Thus, in some cases of ReA we can observe activation of structural alterations. The results show reliable difference in etiological factors, progression, and duration of the disease.

In its turn, analysis of the etiological factor showed that, exactly in the I group (with *Chlamydia trachomatis* etiology) both male and female patients had reliable ( $p<0.02$ ;  $p<0.001$ , respectively) rise COMP in blood serum compared to the control group. At the same time, we determined that, progression of the pathology in these patients was relevant to structural alterations in cartilages. Figure 2 demonstrates that, long-lasting progression of the disease was reliably ( $p<0.01$ ) accompanied by the rise of COMP. It was particularly expressed in women, who had  $3672.8\pm176.6$  ng/mL, which was 2.5 fold ( $p<0.05$ ) higher than in men ( $1421.8\pm412.3$  ng/mL). These data certainly indicate

presence of gender differences in the progression of the disease and probable involvement of sexual hormones in its genesis.

The study of COMP level in blood serum of the patients suffering ReA showed characteristic dynamic changes occurring with the prolongation of the term of disease. Table 3 shows that, in the I group COMP started rising within initial stages of the disease ( $p<0.05$ ) and intensified with the prolongation of the term of the disease. So, on the basis of the obtained results, we can make a conclusion about long-lasting progression and start of early destruction of cartilage in the structure of joint, especially in women, within initial stages of *Chlamydia trachomatis* ReA. Besides that, it can indicate intensification of structural changes in joints and development of secondary osteoarthritis in initial stages of the disease.

Parameters of cartilage COMP in patients with ReA

Groups	COMP, ng/mL		p
	Male	Female	
Control, n=20	814.7±52.2	912.2±112.2	-
1, n=60	2746.2±393.5	3051.2±165.5	p <sub>э</sub> <0.02; p <sub>а</sub> <0.001
2, n=30	1003.7±119.1	1132.5±302.7	p <sub>э</sub> >0.05; p <sub>а</sub> >0.05
3, n=30	987.2±715.2	1089.9±955.7	p <sub>э</sub> >0.05; p <sub>а</sub> >0.05
Total, n=120	1343.2±1101.2	1359.2±1002.8	p <sub>э</sub> >0.05; p <sub>а</sub> >0.05

Note. p<sub>э</sub> – male and p<sub>а</sub> – female reliable parameters compared to the control group.

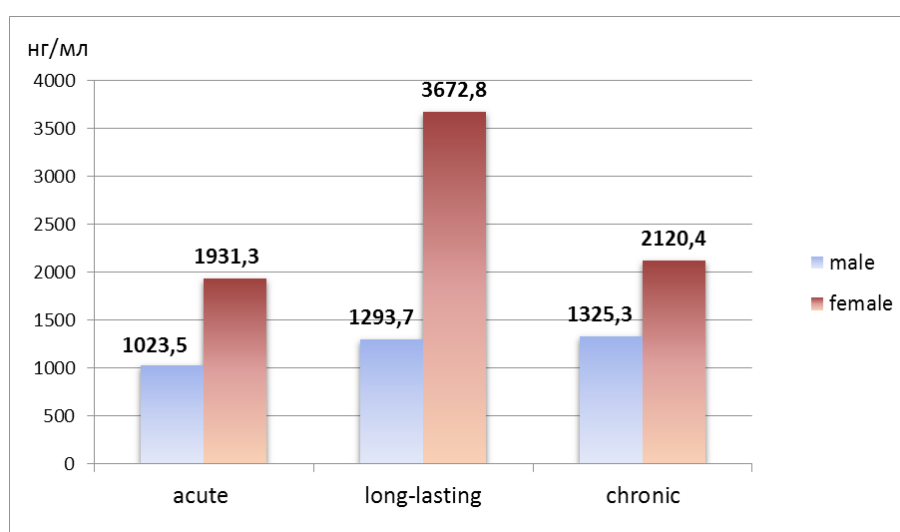


Fig. 2. Change of COMP level in patients with ReA depending on the progression of the disease.

Change of COMP level depending on the term of disease

Term, month	1 group, n=60	2 group, n=30	3 group, n=30	Control, n=20
0-12	1348.1±97.9*	901.7±95.2	843±231.4	865,5±82,2
12-24	2293.5±223.2*	913.3±118.8	909±219.2	
24-36	2870.2±191.3*	1055.8±342.1	891±105.8	
36-48	3012.6±234.2*	1456.5±201.2*	1132±124.3	
More than 48	3144.1±301.5*	2011.4±121.9*	1897±100.8*	

Note. \* – p<0.05 reliability of the data compared to the control.

### Conclusion

1. Thus, secondary osteoarthritis develops within initial stages of ReA and rise of cartilage oligomer matrix protein (COMP) indicates early destruction of cartilage. ReA with Chlamydia trachomatis etiology and long-lasting progression is characterized by intensification of structural alterations in joints.

2. It is worth mentioning, that x-ray images of knee joint of the patients with Rea showed development of secondary osteoarthritis within initial years of the disease in 20% of the patients. Figure 3 demonstrates that, alterations develop with the progression of the disease. In 23.4% of the cases in thirty six months from the start of ReA we could observe III and IV stages of osteoarthritis, in other words there was need of endoprosthetics.

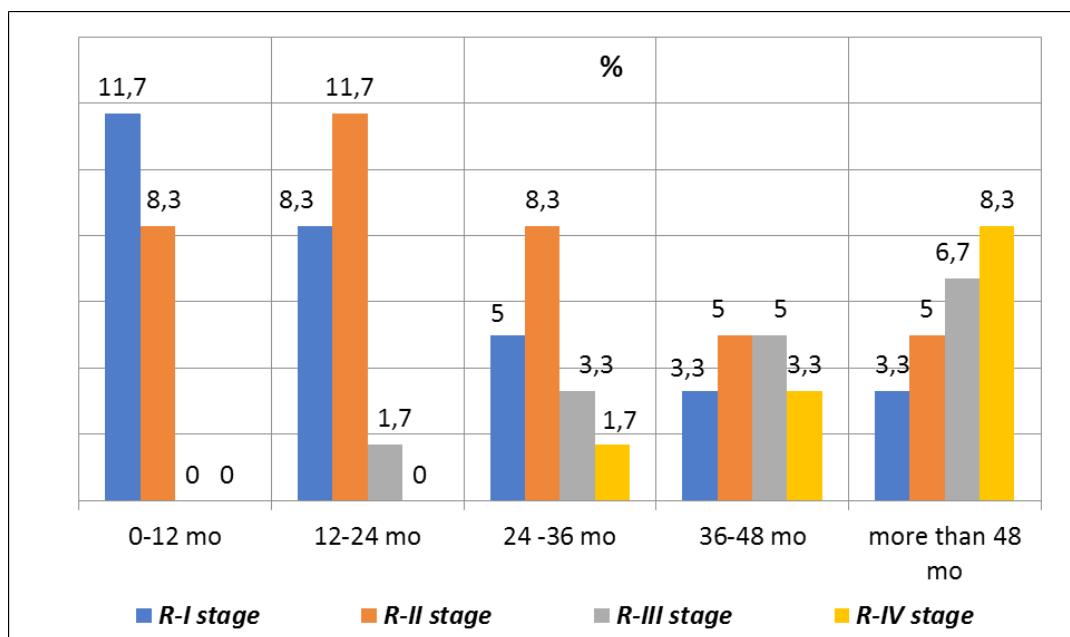


Fig. 3. Development of secondary osteoarthritis in ReA patients of the I group (*Chlamydia trachomatis* etiology).

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#### SPECIFIC DESTRUCTION OF THE JOINT STRUCTURE IN REACTIVE ARTHRITIS

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**Objective:** Study of dynamic changes of cartilage oligomere matrix protein level in patients with reactive arthritis dependent on its clinical presentation for the definition of early cartilage destruction. **Materials and methods:** The study enrolled 120 patients in the age of  $36,9 \pm 12,3$  classified into three groups according to the trigger factors and 20 healthy subjects. **Results:** On the basis of the obtained data about early stages of reactive arthritis with *Chlamydia trachomatis* etiology, long-lasting progression, and particularly rise of cartilage oligomere matrix protein in women we can make a conclusion about the initiation of early cartilage destruction in the structure of a joint. **Conclusion:** Besides that, it indicates activation of alteration in the articulate structure and development of secondary osteoarthritis in early stages of the disease.

**Key words:** reactive arthritis, cartilage oligomere matrix protein, joints syndrome destruction, *Chlamydia trachomatis*.