Methods: We studied the serum from 30 apparently healthy individuals, and 108 rheumatoid arthritis patients. Antibodies to CP were determined by enzyme immunoassay using immobilized granulated antigen preparations (modification by Gontar et al. 2009). Amount of CP was determined by enzyme immunoassay according to the method of I.S. Kuzmina et al. (1991) using commercial diagnostic agent manufactured by Mechnikov Research Institute for Vaccines and Sera.

Results: Enzyme immunoassay showed a mean level of CP antibodies in donor sera of 0.020±0.068 optical density units. The level of normal values of specific antibodies determined as M Abs included an extinction value in the range 0 – 0.086. The mean value of oxidative activity and the amount of CP in healthy people was 716±25.3 and 921±32.6 ng/ml, correspondingly. In the process of study we revealed a reliable increase in CP antibody count, the activity and amount of CP in patients with rheumatoid arthritis while in all cases the parameters under study correlated with the degree of disease activity (p<0.05); at activity degree I CP antibodies were 0.099±0.011; CP activity was 954±48.1; CP content was 1292±73.4. At activity degree II CP antibodies were 0.138±0.007; CP activity was 1163±39.6; CP content was 1368±89.5; CP activity was 1794±102.8. After a course of hospital treatment was completed, we noted a reliable decrease in the activity and amount of CP (at degree I of rheumatoid arthritis activity p<0.01, at degree II of rheumatoid arthritis activity p<0.01 for both parameters; at degree III, p<0.05) compared with baseline findings. A decrease in CP antibodies shows decelerated dynamics, especially in patients with pronounced disease activity, which indicates severe disorders in the immunity that cannot be cured completely within 30 – 40 days of hospital treatment course.

Conclusion: Determination of CP antibodies, as well as oxidative content of CP and its oxidative activity can serve as indicators of the activity of rheumatoid arthritis, as well as an accessory criterion of the effectiveness of administered therapy.

Disclosure of Interests: None declared

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AB0111

FAMILIAL RHEUMATOID ARTHRITIS WITH LATE ONSET (50 YEARS AND OLDER) IN CLINICAL PRACTICE ACCORDING TO THE ALL-RUSSIAN ARTHRITIS REGISTER (OREL)

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Background: Relatives of the 1st degree of relationship (RFDR) of a patient with rheumatoid arthritis (RA) have the higher risk for the development of RA, associated with a 2-5 fold increase.

Objectives: Clinical characteristics of probands with RA onset at an older age (50 yrs and older) and their RFDR, which had RA, other rheumatic diseases (RD) and other autoimmune diseases (AID).

Methods: 2766 RA pts with onset of the disease at the age of 50 yrs and older, residents of Moscow and the Moscow region, were included in the Russian register of arthritis OREL in the period from 01.06.2012 to 31.12.2018. 67 probands with RA (2.4%) had RFDRs with RA, other RDs, other AIDs (86 RFDRs in total). Disease activity (by DAS-28), X-ray stage - (by modified Steinbrocker), functional disorders (by functional class - FC), immunological characteristics (RF, ACPA), etc., characterizing the clinical manifestations of RA were evaluated in accordance with the requirements of the current national working RA classification. RFDRs were divided into groups depending on relationship to proband father, mother and siblings with RA, RFDRs with other RD and RFDR with other AID; serological affiliation to RF and ACPA; association with concordant and discordant RFDR, as well as on number of RFDRs (1 and 2 or more).

Results: 67 probands with RA were investigated: 9 men, 58 women, average age in debut (m ± s) 58.78 ± 6.23 yrs; the average age at the time of the last study was 68.8 ± 6.38 yrs, average duration of RA was 10.14 ± 5.69 yrs. 6 (9%) probands had FC1, 45 (69%) - FC2, 13 (20%) - FC3, 1 (2%) - FC4, 2 - without data; 34 (53%) had high activity, 23 (36%) -medium, 6 (9%) - minimal, 1 (2%) - remission, 3 without data; RF-positive (RF+) - 77.0%, ACPA-positive (ACP+) - 71.7%. 19 probands (29%) were Cohort X-ray stage III; 17 (25%) stage IIb, 10 (15%) stage III, 3 (5%) - stage 4, 8 - without data. 61 (91%) had advanced clinical stage, 6 (9%) - late. Among 86 RFDRs (67 RF-, 19, RF+) 10 had father with RA (8 RF+ probands, 2 RF-), 2 father with other RDs (all RF+), 10 - mother with RA (all RF+), 7 - mother with other RDs (4 RF-, 3 RF+), 26 - siblings with RA (20 RF+, 6 RF-), 11 - siblings with other RDs (9 RF+, 2 RF-), 20 - RD with other RDs (14 RF-, 6 RF+). Probands with concordant RFDRs had more pronounced changes: high activity (66% versus 32%), X-ray erosive stages (67% versus 60%) and stages III and IV (32% versus 6%); FC3 and FC4 (23% versus 12%) and a tendency towards more frequent development of the late clinical stage (11% versus 2%). Comparison of clinical signs in probands with 1 RFDR (49 cases) and probands with 2 or more RFDRs (17 cases) showed a significant predominance of high inflammatory activity (61% versus 41%), erosive arthritis (77% versus 57%), and a tendency to development of a late clinical stage (10% versus 6%) in probands with 1 RFDR. In probands with RF-positive RA, the predominance becomes more significant: high inflammatory activity (69% versus 43%), erosive arthritis (90% versus 54%), late clinical stage (13% versus 7%), and a tendency to severe functional impairment - FC3 and FC4 (22% vs. 15%). ACPA-positive probands showed the same tendency as RF-positive.

Conclusion: The clinical picture of probands with RFDRs with RA, other RDs, other AIDs is generally characterized by more pronounced changes in probands with RA who have RFDRs with concordant diseases in the form of more often high inflammatory activity, less often moderate, erosive arthritis, more frequent development of advanced 3 and 4 X-ray stages (every fifth). Also, more pronounced changes develop in probands with RA who have only 1 RFDR compared to probands with 2 or more RPSRs. In this group, the differences are even more pronounced in the case of RF-positive probands. In cases with ACPA-positive probands, the same trends persist. Careful interview of a proband with RA about the presence of RFDRs with RA, other RDs and other AIDs will make it possible to predict its course more correctly and facilitate planning of rational therapy.

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AB0112

TFN INHIBITOR MONOTHERAPY IN RHEUMATOID ARTHRITIS: IS THERE REALLY A DIFFERENCE IN COMPARISON WITH COMBINATION THERAPY WITH CSDMARDS IN REAL-LIFE?

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Background: In Rheumatoid Arthritis (RA), tumor necrosis factor inhibitors (TNFi) in combination with conventional synthetic disease-modifying antirheumatic drugs (csDMARDs) has shown advantages concerning efficacy and immunogenicity in comparison with monotherapy. However, in clinical practice, up to 40% of patients under biological DMARDs (bDMARDs) are on monotherapy.

Objectives: To compare the efficacy outcomes of TNFi in monotherapy and in combination therapy in a RA multicentric cohort.

Methods: Retrospective, cross-sectional study including all the RA patients under TNFi followed at our Rheumatology Department and registered in the national database. Demographic, clinical and laboratory data and disease activity, disability and other patient-provided data (adherence) were collected at the last visit of 2019 from each patient. Mann-Whitney U and chi-square tests were used to test the comparison analysis between the two groups (continuous and categorical variables, respectively).

Results: A total of 144 patients were included: 84% were females; at the last visit of 2019, the mean age was 56.3±10.9 years and the mean disease duration was 18.3±10.2 years (67% of patients were treated with TNFi at the last visit of 2019). Mann-Whitney U and chi-square tests were used to compare the variables between the two groups (continuous and categorical variables, respectively).

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