Results: Among 96 RA patients, the mean of serum IgG4 was 48.0±45.4 mg/dL and 6.3% had elevated serum IgG4. The mean serum IgG4/IgG ratio of RA patients was 3.5%±2.8% (range 0.2%–16.9%). There was no patient with elevated serum IgG4 in ankylosing spondylitis, systemic lupus erythematosus, Sjögren’s syndrome, and inflammatory myositis. When the patients were divided according to clinical activity, the percentages of the positive serum IgG4 were 25% in active disease group and 4% in low activity group. However, the serum IgG4 levels of the RA patients with active disease activity were not significantly higher than those of the RA patients with low disease activity (58.3±44.3 mg/dL vs. 29.5±20.1 mg/dL). No significant relationship was observed between the ratio of IgG4 total IgG and disease activity. The IgG4 concentrations and total IgG IgG4 ratios were similar between RA and the other autoimmune diseases (p>0.05).

Conclusions: Our results showed that elevated serum IgG4 in RA is relatively common. However the presence of the elevated serum IgG4 was not associated with disease activity of RA. Further investigations are needed to explore the clinical significance in a larger study population.

REFERENCES:

Disclosure of Interest: None declared

AB0213 CORRELATION BETWEEN COMPONENTS OF THE DAS28 SCORE AND HEALTH ASSESSMENT QUESTIONNAIRE IN EARLY RHEUMATOID ARTHRITIS
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Background: The health assessment questionnaire (HAQ) in rheumatoid arthritis (RA) has been widely validated as a patient reported outcome measure (PROM). In the United Kingdom regulatory bodies such as the national institute of health-care and clinical excellence (NICE) have been using improvement in HAQ as a surrogate for efficacy of drugs in RA and has informed their calculation of quality adjusted life year, and calculating health costs using incremental costs effectiveness ratios. Most clinical trials in RA report their primary outcome as improvement in clinical parameters such as swollen and tender joints, inflammatory markers and patient and physician global assessment of disease. What is not clear how the two sets of parameters interact. Some reports1 indicate that there is a strong correlation in early disease, but this has not been validated.

Objectives: We set out to determine the relationship between HAQ scores and clinical parameters of the Disease activity score (DAS28) in addition to physician global.

Methods: Patients were recruited from a single centre from the RAMS study in the North west of England. This is a study of patients with early newly diagnosed RA commencing methotrexate. A subset of patients filled in the HAQ questionnaire and this as used as an outcome variable using linear regression and the swollen joints, tender joints, patient global assessment of disease as well as physician global assessment of disease in addition to inflammatory markers were used as explanatory variables. These were then adjusted for age.

Results: 81 patients were included in the analysis. median age was 63.1 years (IQR 52.9,72.5), 50 (61.7%) were female, the median HAQ score at baseline was 1 (IQR 0.5,1.5) the median DAS28 score 5.3 (IQR 4.5,6.3). Tender joints at baseline correlated well with HAQ score Beta=0.058 95% CI, 0.04,0.08 (p<0.01). Swollen joints did not correlate with the HAQ Beta=0.000 (95%CI −0.3,0.3). Physician global correlated well with disease beta=0.014 (95%CI 0.005,0.022). Patient global assessment correlated well with HAQ (beta 0.014 95% CI 0.008,0.020), CRP did not correlate with HAQ (beta 0.00261 95% CI −0.002,0.007)

Conclusions: In this small study, patient and physician related outcome measures correlate with HAQ scores at baseline more than measures of joint swelling and inflammatory markers. This indicates that using HAQ as an outcome measure underestimates the effect of treatment. When assessing the efficacy of drugs using HAQ this should be taken into account. Validity of the approach needs to be reviewed.


Disclosure of Interest: None declared

AB0214 THE TRAJECTORY OF DISEASE ACTIVITY IN RHEUMATOID ARTHRITIS IN THE FIRST TWO YEARS OF TREATMENT IN AN ASIAN RA COHORT
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Background: Response to disease-modifying antirheumatic drugs (DMARDs) is heterogeneous. Clinical information and baseline characteristics do not allow reliable prediction of which trajectory patients will follow after DMARD initiation.

Objectives: Weanalysed the change in disease activity over the first two years of treatment in rheumatoid arthritis (RA) to identify different treatment response patterns among RA patients initiating DMARDs. We wanted to establish a predictive model for identifying patients with different treatment response patterns.
Methods: We selected patients from our prospective RA disease registry who have been treated for three months or fewer at study entry. We analysed the change of the disease activity, as defined by the DAS28-ESR, over the subsequent two years. A predictive model with parameters from three time points is proposed to stratify patients according to the outcomes.

Abstract AB0214 – Figure 1

Results: We analysed the data from 179 patients over 1044 study visits. We discerned three groups of patients according to disease activity trajectories: the first group (53%) has high DAS at study entry and approach remission after 18 months; the second group (22%) has high DAS at entry that remained elevated throughout the study period; and the third group of patients (25%) started with moderately high DAS and reached remission after 3 months of treatment. Patients at risk of being in the third group can be identified using data from three time points, at initiation of DMARDs, at 3 months and at 6 months.

Conclusions: RA patients showed three distinct disease activity trajectories with treatment. Our model can categorise patients into these groups.

REFERENCES:

Disclosure of Interest: None declared

AB0215

ASSOCIATION OF RHEUMATOID FACTOR IMMUNOGLOBULIN A SEROPOSITIVITY WITH RISK OF EROSIVE ARTHRITIS IN PATIENTS WITH RHEUMATOID ARTHRITIS: AN OBSERVATIONAL STUDY

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Background: In patients with rheumatoid arthritis (RA), erosive arthritis is a major determinant of long-term prognosis. Seropositivity for immunoglobulin (Ig) M rheumatoid factor (RF) or cyclic citrullinated peptide antibodies (anti-CCP) are risk factors for erosive arthritis. However, RA patients can also present IgA RF. The risk for erosive arthritis associated with RF IgA seropositivity is not established.

Objectives: To evaluate the risk for erosive arthritis associated with IgA RF seropositivity in RA patients.

Methods: Cross-sectional observational study, including RA patients (fulfilling the 2010 ACR/EULAR classification criteria) and consecutively observed in a hospital-based rheumatology outpatient clinic, from April to August 2017. At time of inclusion, patient characteristics were evaluated, including: gender, age, RA duration since time of diagnosis, smoking habits, seropositivity for IgA RF, IgM RF and anti-CCP, erosive arthritis in hand and feet X-rays. Risk association for erosive arthritis was analysed with univariate and multivariable logistic regression models for the putative risk factors and confounders. Odds ratios (OR) and 95% confidence intervals (CI) of IgM RF, IgA RF, and anti-CCP seropositivity for erosive arthritis were estimated. Statistical significance was set at 0.05.

Results: 86 patients were included. The univariate logistic regression showed significant positive associations of IgA RF, IgM RF and anti-CCP with erosive arthritis. In the multivariate analyses, adjusting for confounders (gender, age, disease duration and smoking), the OR for erosive arthritis associated with IgA RF, IgM RF and anti-CCP were respectively: OR=2.42 (95% CI 0.72–8.07; p=0.152); OR=3.54 (95% CI 1.16–10.83; p<0.05); OR=4.13 (95% CI 1.33–12.82; p<0.05). The seropositivity for IgM RF, IgA RF and anti-CCP were strongly associated among each other (Chi-square test with p<0.001 for all associations).

Conclusions: In this RA cohort, the IgA RF was associated with erosive arthritis in univariate analysis, but did not prove to be an independent risk factor in multivariate regression, due to its strong association with IgM RF and anti-CCP. Determination of IgA RF does not seem to add predictive value for erosive arthritis in RA patients.

Disclosure of Interest: None declared

AB0216

LOW MORTALITY RATE IN ITALIAN RHEUMATOID ARTHRITIS PATIENTS FROM A TERTIARY CENTRE. PUTATIVE IMPLICATION OF A LOW ANTICARBAMYLATED PROTEIN ANTIBODIES PREVALENCE

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Background: Rheumatoid Arthritis (RA) is a chronic systemic inflammatory disorder associated with increased mortality, in particular from cardiovascular (CV) disease, infections and cancer. We recently demonstrated a incidence mortality rate (IMR) in 654 RA patients enrolled over a 6 year period in a South-Italian tertiary Rheumatology Centre lower than that reported in the Norfolk Arthritis Registry.

Objectives: The present study is devoted to investigate differences in IMR between our series and other European tertiary centre cohorts. Furthermore we evaluated the role, if any, of Anticarbamylated protein antibodies (anti-CarP Ab) in modulating the low IMR detected in our patients.

Methods: Clinical charts of patients consecutively admitted to our centre, from January 1st, 2008 to December 31st, 2014 were reviewed. IMRs and causes of death as assessed at December 31st 2015, were registered. Sera collected at the time of admission to our centre in 61 patients representative of our RA cohort were investigated for the presence and the level of anti-CarP Ab. Demographic and clinical features, mortality rates and prevalence of anti-CarP Ab detected in our series were compared with those reported in the Better Anti-rheumatic Farmaco-therapy (BARFOT) cohort, the Leiden Early Arthritis Clinic cohort (Leiden EAC) and a Spanish cohort.

Results: Six hundred and eight patients were observed for a median of 3.51 years. All causes and cause-specific IMRs were significantly lower in our cohort with respect to the BARFOT and the Spanish cohort, while only all causes and CV IMRs were significantly lower in our series with respect to the Leiden EAC. These discrepancies might depend on demographic and clinical differences among the various cohorts. Nevertheless, we found to fail putative differences with respect to each North European cohort, but we detected a significantly lower prevalence of anti-CarP Ab in our series with respect to that reported in the other European cohorts considered (table 1).

Conclusions: In conclusion, we confirm that the mortality rate in our South Italian RA cohort is lower than that detected in patients from both North and South European countries.

We detected a very low prevalence of anti-CarP Ab in our sample representative of the entire cohort. Whether this is the aspect underpinning the low mortality rate detected in our series, awaits to be furtherly investigated.