between YORA and LORA were not significantly different at initial presentation (table 1). The ultrasound characteristics differed between these two groups. LORA patients were more likely to have shoulder biceps tendon tenosynovitis (GS; p=0.026, PD; 0.037), elbow joint synovitis (GS; p=0.010, PD; p=0.037), MCP1 (PD; p=0.032) and MCP5 (GS; p=0.035) synovitis, compared to YORA patient. YORA patients were more likely to have MTP synovitis (GS; p=0.013) compared to LORA patients.

**Conclusions:** This is the first study to describe the difference of both clinical and sonographic inflammation of YORA and LORA in recent-onset DMARD-naïve RA patients in a longitudinal study. There are differences in US-detected joint and tendon inflammation despite similarities in clinical characteristics. The prognostic value of the differences in US pathology between these two groups should be further explored.

**Disclosure of Interest:** None declared

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**OP01554**

**PROSPECTIVE OBSERVATIONAL STUDY TO EVALUATE THE USE OF MUSCULOSKELETAL ULTRASOUND TO IMPROVE RHEUMATOID ARTHRITIS MANAGEMENT: INTERIM ANALYSIS OF THE ECHO STUDY**

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**Background:** Musculoskeletal Ultrasound (MSUS) has been shown to be superior to clinical examination in the detection of synovitis in patients with Rheumatoid Arthritis (RA), and can be used to improve diagnostic accuracy and potentially monitor disease changes in order to make treatment decisions aimed at optimising patient care. Since the creation of the Canadian Rheumatology Ultrasonography Society (CRUS) in 2009, an increasing number of rheumatologists has been trained in the use of MSUS.

**Objectives:** The overall study objective is to compare the effectiveness of MSUS to Routine Care (RC) as a disease management tool in patients with moderate-to-severe RA for whom a change in treatment is indicated. In addition, the predictive power of MSUS assessments has been assessed here.

**Methods:** Echo is a prospective two-cohort, quasi-experimental study of patients diagnosed with active moderate-to-severe RA managed either with MSUS (within CRUS) or as per RC. To be eligible for the study patients must require a change in treatment as per the judgment of the treating physician. Patients are followed for 1 year with assessments at baseline, 3, 6, 9, and 12 months. Key outcome measures of interest include CDAI LDA/Remission, DAS-28 LDA/Remission, patient satisfaction (TSQM) and patient perception of participation in disease management (PAM-13).

**Results:** A total of 383 patients (71.5% female) with a mean (SD) age of 58.7 (11.7) years and disease duration of 7.0 (10.0) years were enrolled, without any significant differences between treatment groups. At baseline, a greater proportion of patients in the MSUS group were treated with a biologic DMARD (bDMARD; 50.3% vs 35.8%, p=0.004); patients in the RC group were more likely to be treated with a non-biologic DMARD (nbDMARD; 84.2% vs 91.5%, p=0.027). Over time, a comparable proportion of patients in the two groups started/switched a bDMARD (21.6% vs 15.6%, p=0.126) or added/switched a nbDMARD (18.7% vs 23.6%, p=0.248). The overall number of treatment modifications was also similar between groups (3.0 vs 2.7, p=0.236).

Upon adjusting for age, gender, previous bDMARD treatment, and baseline parameter level, no differences between the two treatment groups with respect to CDAI LDA/Remission, DAS-28 LDA/Remission, and TSQM score were observed during follow-up. However, the PAM-13 score was significantly higher in the MSUS group (69.6 vs 64.2, p=0.02).

In the MSUS group, higher total US erosion score at baseline was associated with a lower rate of CDAI LDA at 12 months (OR=0.86; p=0.047); higher total PD synovitis score at baseline was associated with a lower rate of CDAI LDA at 6 months (OR=0.90; p=0.010); and, higher total synovitis GREY scale at baseline was associated with lower rates of DAS28 LDA (OR=0.93; p=0.026) and DAS28 Remission (OR=0.94; p=0.061) at 6 months.

**Conclusions:** MSUS assessments can be useful predictors of future disease remission in patients with RA. MSUS may be associated with increased patient perception of participation in disease management and patient activation.

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**OP0155**

**ULTRASOUND AS AN OUTCOME MEASUREMENT TOOL FOR OPTIMISED MONITORING OF GOUT. VALIDATION OF THE OMERACT ULTRASOUND DEFINITIONS OF GOUT ELEMENTARY LESIONS**

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**Objectives:** To evaluate ultrasound (US) as an outcome measurement instrument for monitoring gout patients during urate lowering therapy using the OMERACT US Working Group’s 2015 definitions of US elementary lesions in gout.

**Methods:** US examination (28 joints, 26 tendons) were performed in patients with microscopically verified gout who either initiated or increased urate lowering therapy. Joints and tendons were evaluated for the four OMERACT elementary lesions of gout (Double contour, Tophus, Aggregates and Erosions). Furthermore, subcutaneous (SC) oedema was registered and synovitis was graded by grey scale (GS) and colour Doppler (CD) (both graded 0–3). A sum score was