The same upward trend was found with regard to disease duration before arthroplasty: a significant difference between group A and group B (16.98 y vs 21.66 y; p = 0.03) and between group A and group C (16.98 y vs 22.93 y; p = 0.00).

The rate of implant survival at 5, 10 and 15 years were comparable (from 84% to 89%); whereas 50% of eligible implants lasted 20 years or more (Figure 2).

**Figure 2.** Kaplan-Meier survival curve of implants.

The year of surgery was found to be significantly related to implant survival [Hazard Ratio (HR) 1.001, confidence interval (CI) 1.0001-1.0006; p < 0.001] as well as the presence of complications (HR 3.69, CI 1.82-7.48; p < 0.001) in multivariante analysis. Furthermore, prostheses of polyarticular RF-neg patients had more possibilities to last longer than those of S-JIA patients (HR 0.23, CI 0.09-0.53; p = 0.00) as well as implants of all polyarticular RA (RF-pos and neg together) (p < 0.001).

**Conclusion:** We observed an upward trend of both age at arthroplasty and disease duration before the first arthroplasty over time. JIA category, year of implants and the presence of complications significantly affected implant survival. Future researches should assess functional outcome and survival of implants according to medical therapy and different surgical approaches.

**Disclosure of Interests:** None declared

**DOI:** 10.1136/annrheumdis-2020-eular.1747

**THU0505 MUSCULOSKELETAL ULTRASOUND MONITORING DURING MTX TAPERING IN JIA: A PROSPECTIVE BLINDED COHORT STUDY**

F. Licciardi1, M. Dellepiane1, C. Covizzi2, F. Figus2, I. Azzolin2, D. Montin1, A. Iagnocco1, R. Margherita Children Hospital, Rheumatology Department, Turin, Italy; 2Università degli Studi di Torino, Turin, Italy

**Background:** Juvenile idiopathic arthritis (JIA) is the most common chronic rheumatic disease in children. Musculoskeletal ultrasound (MSUS) is a reliable tool in the assessment of chronic inflammatory arthropathies. MSUS in JIA has demonstrated a higher sensitivity for detecting synovitis and tenosynovitis as compared to physical examination. The examination of subclinical synovitis (Sub-S: MSUS+/ physical examination -) seems more frequent in wrist and foot joints; the clinical significance of Sub-S in real-life practice is still debated. Methotrexate (MTX) is the most widely used first-line DMARD in JIA therapy. Weekly treatment with MTX leads to clinical remission (CR) in 50-70% of patients. After a variable period of CR (usually 6-18 months), MTX is discontinued. Relapse rate after MTX suspension ranges between 40-50%; no predictors of disease flare have been identified so far.

**Objectives:** We designed a cohort study in order to explore if MSUS monitoring during MTX tapering was able to predict disease flare.

**Methods:** JIA patients in CR (as defined by the JADAS score) for at least 12 months were enrolled in the study. Patients at first attempt of suspension (G1) were tapered as follows: 1 week of suspension every 3 weeks + 1 dose every 2 weeks for 3 months; if CR persisted, MTX was stopped. Patients who had a previous flare during/after MTX tapering (G2) had a similar tapering schedule but the step with 1 MTX dose every 2 weeks lasted 6 months. All patients underwent a complete MSUS of 48 joints every 3 months; clinicians who performed physical examinations and follow-up were blinded to US findings for the entire study period.

**Results:** 18 consecutive patients were enrolled between April 2018 and September 2019; patients had prevalently oligoJIA (55.5%) and RF- polyJIA (22.2%). Patients had been treated with MTX for 24.7 months (17.7-48.3). CR had been achieved 4.2 months after MTX start; 61.1% were at their first attempt of MTX tapering (G1). Baseline MSUS: at T0 MSUS detected 9/18 patients (50.0%) with Sub-S (MSUS+); Affected sites at T0 were distributed as follows: 4 MCP joints, 9 MTP joints, 1 H-P joints, 11 knees. No significant differences resulted in comparing demographic and baseline disease features between MSUS- and MSUS+ patients at T0. Follow-up MSUS: 14 patients (77.8%) completed the entire study protocol, 4 patients are still ongoing. 7 patients relapsed: 42.9% during tapering, 1 of them relapsed during a VZV infection and was excluded from further analysis. We considered as Tlast-MSUS the last available MSUS before relapse or final MSUS (i.e. three months after MTX withdrawal) for not-relapsed subjects. At Tlast patients had at least 1 Sub-S. Sub-S per patient at Tlast were more than Sub-S at T0 (2.85 vs 0.53, p=0.03) but the presence of Sub-S was not related with disease flare (50.0 vs 44.4%, p=1). MSUS found 27 Sub-S of the small joints (sMSUS): 88.9% were in the feet, they had an OMERACT grading of 1. sMSUS+ patients were older (8.7 vs 3.9, p=0.002) therefore a weight-induced sub-S not related with JIA could be presumed.

Kaplan-Meier curves were analyzed comparing MSUS results at T0 and Tlast, both considering all Sub-S and excluding small feet joints (pMSUS). The best performance was achieved with MSUS at Tlast and pMSUS (figure below, p=0.11).

**Conclusion:** • Sub-S are present in 50% of patients in clinical remission >12 months.
• Sub-S in older patients interest often feet small joints; these Sub-S may be of mechanical origin and are not associated with disease flare.
• Sub-S increase during MTX tapering.

**References:**


**Disclosure of Interests:** Francesco Licciardi: None declared, Marta Dellepiane: None declared, Carlotta Covizzi: None declared, Fabiana Figus: None declared, Irene Azzolin: None declared, Davide Montin: Speakers bureau: Not relevant for the topic, Annamaria Iagnocco Grant/research support from: Abbvie, MSD and Alfasigma, Consultant of: AbbVie, Abiogen, Alfasigma, Biogen, BMS, Celgene, Eli-Lilly, Janssen, MSD, Novartis, Sanofi and Sanofi Genzyme, Speakers bureau: AbbVie, Alfasigma, BMS, Eli-Lilly, Janssen, MSD, Novartis, Sanofi

**DOI:** 10.1136/annrheumdis-2020-eular.4151

**THU0506 LONG-TERM EFFECTIVENESS AND SAFETY OF CANAKINUMAB AS A SECOND BIOLOGIC AFTER TOCILIZUMAB IN CHILDREN WITH EARLY AND LATE JIA WITH ACTIVE SYSTEMIC FEATURES**

A. Alexeeva1,2, E. Krekhova1, T. Dvoryakovskaya1, K. Isakova1, A. Chomakhidze1, E. Chistyakova1,2, O. Lomakina1, R. Denisova1, A. Mamutova1, A. Fetisova1, M. Gautier1, D. Vankova1, M. Shingarova2, A. Ailtsevkaya2, A. Moskaleva1, I. Krulin1,2, E. Alexeeva1,2, E. Krekhova1, T. Dvoryakovskaya1, K. Isakova1

1Regina Margherita Children Hospital, Rheumatology Department, Turin, Italy; 2Università degli Studi di Torino, Turin, Italy

**Background:** Canakinumab (CAN) is often used as second biologics in juvenile idiopathic arthritis with active systemic features (sJIA). However, there are little information about its long-term efficacy and safety.

**Objectives:** To evaluate the long-term effectiveness and safety of CAN in juvenile idiopathic arthritis with active systemic features (sJIA).

**Methods:** Canakinumab (CAN) was administered as second biologics after tocilizumab (TOC) in sJIA patients depending on the duration of the disease.

**Results:** 491

**Conclusion:** CAN is a safe and effective treatment for sJIA patients in clinical remission >12 months.

**Disclosure of Interests:** None declared

**DOI:** 10.1136/annrheumdis-2020-eular.5706

Thursday, 04 June 2020

© 2020 BMJ Publishing Group Ltd

All rights reserved. No part of this publication may be reproduced, stored in any form or transmitted in any means without the prior permission in writing from the publisher.
ASSOCIATION BETWEEN JUVENILE IDIOPATHIC ARTHRITIS AND AUTISM

R. Beesley1, 1Juvenile Arthritis Research, Tonbridge, United Kingdom

Background: Juvenile Idiopathic Arthritis (JIA) is a heterogenous group of auto-immune disorders characterised by chronic joint inflammation, diagnosed in around 1 in 1,000 children and young people (CYP) under the age of 16. Autistic Spectrum Condition (ASC) is a neurodevelopmental condition characterised by differences in social communication and sensory perception, as well as restricted interests and repetitive behaviours. Recent estimates from the Centers for Disease Control and Prevention (CDC) suggest that 1.68% of CYP are diagnosed with ASC, with males being more likely to be diagnosed (sex ratio of 4:1) [1]. The causes of both JIA and ASC are complex interactions between genetic and environmental factors. There appears to be some evidence that ASC can be associated with certain parental autoimmune conditions [2], although research into any association between JIA and ASC is sparse with the exception of a review of clinical database information [3].

Objectives: In this parent-led study, the association between JIA and ASC was explored in order to determine if children with JIA, or children who do not themselves have JIA but have at least one first-degree relative with JIA (FDR), are more likely to be diagnosed with ASC.

Methods: Parents of CYP with JIA were invited to complete an online survey, giving details of each of their family including diagnosis status for JIA and ASC, and age of diagnoses. A total of 247 responses were collected, representing 558 CYP. Overall, 202 CYP were diagnosed with JIA from 197 families. The eldest child with JIA from each family was selected (total 197, 66 male and 131 female) and the rate of ASC was compared against the general population using Fisher’s exact tests.

Results: Children with JIA themselves and FDR children were significantly more likely to be diagnosed with ASC.

<table>
<thead>
<tr>
<th>Group</th>
<th>Odds Ratio (95% CI)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>JIA children overall</td>
<td>6.107 (1.760, 21.190)</td>
<td>0.0020  **</td>
</tr>
<tr>
<td>FDR children overall</td>
<td>7.009 (2.033, 24.160)</td>
<td>0.0006  ***</td>
</tr>
</tbody>
</table>

Conclusion: There is an association between JIA and ASC in this study. The majority of children sampled were from the United Kingdom and Ireland; however, we chose to utilise the most recent CDC estimates for ASC prevalence, which are the most recent estimates from the UK were from 2006 and longitudinal data suggests that ASC prevalence continues to increase, likely due to changes in diagnostic criteria and improved recognition of the condition. When using the UK prevalence estimates, JIA children and FDR children remain significantly more likely to be diagnosed with ASC than the general population as a whole.

Future research should focus on confirming these findings in larger, population-based samples.

Disclosures of Interests: None declared

DOI: 10.1136/annrheumdis-2020-eular.5706

LARGE VESSEL VASCULITIS IN A COHORT OF CHILDREN WITH RESISTANT KAWASAKI DISEASE IN SINGAPORE

L. Das1, J. H. T. Tan1, T. Arkachaisri1,2, 1KK Women’s and Children’s Hospital, Rheumatology & Immunology, Singapore, Singapore; 2Duke-NUS Medical School, Singapore, Singapore

Background: Kawasaki Disease (KD) is one of the most common systemic vasculitides in children today. IVIG is the mainstay of treatment, however, about 1/5 of patients do not respond resulting in an increased risk of Coronary Artery Abnormalities (CAA) [1].

Methods: Thirty-one patients were enrolled in this study: the group of early sJIA (with duration shorter than 2 years, 19 patients) and the group of late sJIA (with duration longer than 2 years, 12 patients). At the baseline, information was collected on the characteristics of the onset of the disease, previous therapy and its success. At each visit at least 1 time per year clinical and laboratory characteristics of sJIA severity were assessed. Response to therapy was assessed using the ACRPsfiPed 30/50/70/90 criteria and the SwallQE criteria for inactive disease (WID) and clinical remission.

Results: The most common reason for withdrawal of previous TOC was secondary ineffectiveness (22 cases, 71%); in 6 cases (19%) an allergic reaction was observed; in two cases (6.5%) primary non-effectiveness appeared; and in one case (3.2%) there was marked infusion reaction.

At CAN initiation, sJIA activity was as follows: 15 (12.23) for JADAS-71; 45 (36.5: 72) and 58 (45: 81) for physician’s and patient’s global assessment VAS; and 0.25 (0.62) for the CHAQ disability index. After 12-month treatment, 22 (71%) patients reached WID; 21 on CAN therapy and 1 – after CAN withdrawal due to administrative reason and stable WID. ACR50/70/90 response was achieved by 84.2%/84.2%/64.7% patients in early arthritis group and in 83.3%/75%/75% patients in late arthritis group (p=0.792).

However, 42.1% of patients with early sJIA achieved remission in the first 1.5 years without any further relapse during all the studied period and only 16.7% of patients with late arthritis (p=0.239). In multivariable analysis, it was found that age of sJIA onset (OR (2.5-97.5 CI) 0.353 (0.13 - 0.72), p=0.015), number of joint involvement and low disease activity are predictors of rapid and stable remission.

However, 42.1% of patients with early sJIA achieved remission in the first 1.5 years without any further relapse during all the studied period and only 16.7% of patients with late arthritis (p=0.239). In multivariable analysis, it was found that age of sJIA onset (OR (2.5-97.5 CI) 0.353 (0.13 - 0.72), p=0.015), number of joint involvement and low disease activity are predictors of rapid and stable remission.

A total of 247 responses were collated, representing 558 CYP. Overall, 202 CYP were diagnosed with JIA from 197 families. The eldest child with JIA from each family was selected (total 197, 66 male and 131 female) and the rate of ASC was compared against the general population using Fisher’s exact tests.

Results: Children with JIA themselves and FDR children were significantly more likely to be diagnosed with ASC.

<table>
<thead>
<tr>
<th>Group</th>
<th>Odds Ratio (95% CI)</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>JIA children overall</td>
<td>6.107 (1.760, 21.190)</td>
<td>0.0020  **</td>
</tr>
<tr>
<td>FDR children overall</td>
<td>7.009 (2.033, 24.160)</td>
<td>0.0006  ***</td>
</tr>
</tbody>
</table>

Conclusion: There is an association between JIA and ASC in this study. The majority of children sampled were from the United Kingdom and Ireland; however, we chose to utilise the most recent CDC estimates for ASC prevalence, which are the most recent estimates from the UK were from 2006 and longitudinal data suggests that ASC prevalence continues to increase, likely due to changes in diagnostic criteria and improved recognition of the condition. When using the UK prevalence estimates, JIA children and FDR children remain significantly more likely to be diagnosed with ASC than the general population as a whole.

Future research should focus on confirming these findings in larger, population-based samples.

Disclosures of Interests: None declared

DOI: 10.1136/annrheumdis-2020-eular.876