2019 American College of Rheumatology/Arthritis Foundation Guideline for the Treatment of Juvenile Idiopathic Arthritis: Therapeutic Approaches for Non-Systemic Polyarthritis, Sacroiliitis, and Enthesitis

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Objective. To develop treatment recommendations for children with juvenile idiopathic arthritis manifesting as non-systemic polyarthritis, sacroiliitis, or enthesitis.

Methods. The Patient/Population, Intervention, Comparison, and Outcomes (PICO) questions were developed and refined by members of the guideline development teams. A systematic review was conducted to compile evidence for the benefits and harms associated with treatments for these conditions. GRADE (Grading of Recommendations Assessment, Development and Evaluation) methodology was used to rate the quality of evidence. A group consensus process was conducted among the Voting Panel to generate the final recommendations and grade their strength. A Parent and Patient Panel used a similar consensus approach to provide patient/caregiver preferences for key questions.

Results. Thirty-nine recommendations were developed (8 strong and 31 conditional). The quality of supporting evidence was very low or low for 90% of the recommendations. Recommendations are provided for the use of nonsteroidal antiinflammatory drugs, disease-modifying antirheumatic drugs, biologics, and intraarticular and oral glucocorticoids. Recommendations for the use of physical and occupational therapy are also provided. Specific

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recommendations for polyarthritis address general medication use, initial and subsequent treatment, and adjunctive therapies. Good disease control, with therapeutic escalation to achieve low disease activity, was recommended. The sacroiliitis and enthesitis recommendations primarily address initial therapy and adjunctive therapies.

**Conclusion.** This guideline provides direction for clinicians, caregivers, and patients making treatment decisions. Clinicians, caregivers, and patients should use a shared decision-making process that accounts for patients' values, preferences, and comorbidities. These recommendations should not be used to limit or deny access to therapies.

**INTRODUCTION**

Juvenile arthritis is one of the most common chronic diseases of childhood, with an estimated prevalence of 1 per 1,000 children (1–3). The term juvenile idiopathic arthritis (JIA) defines a heterogeneous collection of inflammatory arthritides of unknown etiology with onset prior to age 16 years and a minimum duration of 6 weeks, following the exclusion of other known causes of synovitis (4). Current International League of Associations for Rheumatology (ILAR) classification criteria divide JIA into 7 mutually exclusive categories defined by the number of joints involved, presence or absence of extraarticular manifestations, and presence or absence of additional markers including rheumatoid factor (RF) and HLA-B27 (4).

All forms of JIA are associated with decreased health-related quality of life and risk of permanent joint damage, and the disease may persist into adulthood, causing ongoing significant morbidity and impaired quality of life (5–13). A number of treatments are available, including nonsteroidal antiinflammatory drugs (NSAIDs), systemic and intraarticular glucocorticoids, and nonbiologic and biologic disease-modifying antirheumatic drugs (DMARDs). Prompt initiation of appropriate therapy is of critical importance in preventing permanent damage and improving outcomes. While earlier disease recognition and expanded treatment options have made good disease control a possibility for many patients, they have also made the decision-making regarding treatments more complex for physicians, caregivers, and patients.

The American College of Rheumatology (ACR) published initial recommendations for JIA in 2011 that provided guidance for the treatment of JIA and for the monitoring of select medical therapies, and an update in 2013 focused on the treatment of systemic arthritis (14,15). The ACR has subsequently transitioned from the RAND/UCLA Appropriateness Method used to generate these prior recommendations to the Grading of Recommendations Assessment, Development and Evaluation (GRADE) methodology, which has the advantages of a more transparent decision-making process and well-defined criteria for moving from evidence to recommendation, including balancing benefits and harms and consideration of patient values and preferences while maintaining methodologic rigor (16).

The goal of this guideline project was to provide updated recommendations for juvenile non-systemic polyarthritis, sacroiliitis, and enthesitis, incorporating recently published data and utilizing the GRADE methodology. Recommendations for the treatment of chronic and acute JIA-associated uveitis were developed concomitantly and are presented separately (17).

**METHODS**

**Methodology overview.** This guideline followed the ACR guideline development process (http://www.rheumatology.org/Practice-Quality/Clinical-Support/Clinical-Practice-Guidelines). This process includes using the GRADE methodology (www.gradeworkinggroup.org) to rate the quality of the available evidence and to develop the recommendations (18–20). ACR policy guided disclosures and the management of conflicts of interest (participant disclosures are available at https://www.rheumatology.org/Portals/0/Files/JIA-Guideline-Disclosures.pdf). Supplementary Appendix 1, available on the Arthritis Care & Research web site

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Guideline development teams. This work involved 5 teams: 1) a Core Leadership Team, consisting of 4 pediatric rheumatologists, who supervised and coordinated the project and assisted with developing the scope of the project and initial Patient/Population, Intervention, Comparison, and Outcomes (PICO) questions and drafting the manuscript; 2) a Literature Review Team, led by an experienced literature review consultant, which completed the literature search and data abstraction and rated the quality of evidence; 3) an Expert Panel, composed of 9 pediatric rheumatologists, who assisted with developing the scope of the project and drafting and refining the PICO questions; 4) a Voting Panel, consisting of 15 pediatric rheumatologists and 2 adult patients with JIA, who assisted with developing the scope of the project and refining the PICO questions and voted on the recommendations; and 5) a Parent and Patient Panel, consisting of 9 adult patients with JIA and 2 parents of children with JIA, who reviewed the collated evidence and provided input on their values and preferences within the context of a separate meeting. Supplementary Appendix 2 (available on the Arthritis Care & Research web site at http://onlinelibrary.wiley.com/doi/10.1002/acr.23870/abstract) presents rosters of the team and panel members. In accordance with ACR policy, the principal investigators and the literature review consultant were free of potential conflicts of interest, and in all teams, >50% of members were free of potential conflicts of interest.

PICO question development and importance of outcomes. The Core Leadership Team drafted the initial project scope, key principles, and examples of relevant PICO questions. The following topics were proposed to the guideline development groups for consideration: acute and chronic anterior uveitis, oligoarthritis, polyarthritis, systemic arthritis, sacroiliitis, enthesitis, and temporomandibular joint arthritis. PICO questions for each topic were developed and discussed at a face-to-face meeting during which the topics were refined. The project scope was subsequently limited to patients with non-systemic polyarthritis, sacroiliitis, and enthesitis, because these were deemed to be the most high-impact areas. The PICO questions for these topics were subsequently reviewed and further refined by the Expert and Voting Panels via e-mail.

Populations (Table 1). While the current ILAR classification criteria have been useful for identifying homogeneous groups of patients for research, more recent data suggest that these categories may not entirely reflect the underlying genetic and clinical heterogeneity of the disease or be relevant for guiding treatment decisions (20–22). For this reason, it was decided to base the current guideline on broad clinical phenotypes rather than ILAR categories, similar to the approach used for development of the 2011 guideline (15). The patient populations addressed in this guideline are defined below. The current recommendations are intended to address typical

<table>
<thead>
<tr>
<th>Table 1. Terms and definitions*</th>
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<tr>
<td><strong>Term</strong></td>
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<tr>
<td>Polyaarthritic population</td>
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<tr>
<td>Risk factors</td>
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<tr>
<td>Moderate/high disease activity</td>
</tr>
<tr>
<td>Low disease activity</td>
</tr>
<tr>
<td>Sacroiliitis population</td>
</tr>
<tr>
<td>Active sacroiliitis</td>
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<tr>
<td>Enthesitis population</td>
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<tr>
<td>Active enthesitis</td>
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</table>

* Disease activity (moderate/high and low) as defined by the clinical Juvenile Disease Activity Score based on 10 joints (cJADAS-10) is provided as a general parameter and should be interpreted within the clinical context. JIA = juvenile idiopathic arthritis. ILAR = International League of Associations for Rheumatology.
patients with the phenotype and may not be applicable to patients with uncommon features or highly refractory disease.

**Polyarthritis.** This group includes children with JIA and polyarthritis (≥5 joints ever involved) and may include children from different ILAR JIA categories but excludes children with systemic arthritis or sacroilitis. These guidelines are not intended to be applicable to children with associated extrarticular manifestations (e.g., psoriasis, uveitis, inflammatory bowel disease) that may also influence treatment decisions. Given the heterogeneity of patients with JIA and polyarthritis, the Expert and Voting Panels initially categorized patients into treatment groups, using combinations of the following categories: 1) presence or absence of risk factors for disease severity and potentially a more refractory disease course, and 2) low disease activity versus moderate/high disease activity.

Risk factors were defined as the presence of one or more of the following: positive anti–cyclic citrullinated peptide antibodies, or joint damage. The Juvenile Arthritis Disease Activity Score (JADAS) was proposed as a means of categorizing disease activity, with the acknowledgment that a number of versions exist, and that validation is not fully complete and cutoff scores may change (23–27). A joint with inactive disease was defined using the ACR definition: presence of swelling (not due to currently inactive synovitis or to bony enlargement) or, if swelling is not present, limitation of motion accompanied by pain, tenderness, or both (28,29). The Voting Panel used the clinical JADAS based on 10 joints (cJADAS-10) and a cutoff of ≤2.5 versus >2.5 to define low versus moderate/high disease activity. Low disease activity was further defined as a cJADAS-10 of ≤2.5 and ≥1 joint with active disease to ensure that active arthritis was also present. Moderate and high disease activity were considered together, because it was thought that treatment approaches would be similar (30). The cJADAS-10 is a sum of the total active joint count (to a maximum of 10), physician’s global assessment of disease activity (0–10 scale), and parent/patient’s global assessment of well-being (0–10 scale). It was also acknowledged that one of the limitations of the JADAS is the lack of standardization of the physician’s and parent’s global assessments. It is therefore recommended that the JADAS be interpreted within the context of the clinical presentation rather than considered an absolute determinant of disease activity.

Because ultimately few data were available to support different treatment approaches based on the risk factors and disease activity categories, patients were often grouped together to provide a recommendation. The few instances in which the Voting Panel made differing recommendations based on disease activity or risk factors are explicitly noted.

**Sacroilitis.** This group includes patients with active sacroilitis who will most likely be classified within the ILAR categories of enthesitis-related arthritis, psoriatic arthritis, and undifferentiated arthritis but may include patients in any of the ILAR JIA categories. In addition to active sacroilitis, patients may or may not have active peripheral joint disease and/or enthesitis to be included in this population. It is anticipated that patients with peripheral spondyloarthritis and no sacroilitis would be treated according to the polyarthritis recommendations included in this update or existing JIA oligoarthritis treatment recommendations from the 2011 ACR JIA guideline (15), depending on the numbers of joints involved. For the purposes of this guideline, patients were considered to have active sacroilitis if they had prior or current magnetic resonance imaging findings consistent with sacroilitis along with clinical examination findings consistent with sacroilitis (e.g., pain with direct palpation of the sacroiliac joints) and/or patient-reported symptoms of inflammatory back pain.

**Interventions.** The pharmacologic and nonpharmacologic therapies considered are listed in Table 2. Both intraarticular and oral glucocorticoids were considered. For the purposes of the recommendations, a bridging course of oral glucocorticoids was defined as a short course (<3 months) of oral glucocorticoids intended to control disease activity quickly during escalation of DMARD or biologic therapy, using the shortest possible duration and the lowest dose needed to control symptoms. The duration of bridging therapy would likely be primarily determined by the anticipated timing of onset of action of the other DMARD or biologic treatment(s). An optimal trial of methotrexate was considered to be 3 months; however, if no or minimal response was observed after 6–8 weeks, it was agreed that changing or adding therapy may be appropriate.

**Outcomes.** Outcomes were selected during the initial face-to-face scoping meeting and subsequently refined by online vote (Supplementary Appendix 3, available on the Arthritis Care & Research web site at http://onlinelibrary.wiley.com/doi/10.1002/acr.23870/abstract). Critical outcomes included disease activity, quality of life, joint damage, and serious adverse events. Pain was selected as an important outcome. While each of these outcomes was thought to be important in decision-making by the guideline development teams, they were not routinely reported across stud-
Table 2. Interventions included in the literature review*

<table>
<thead>
<tr>
<th>Intervention</th>
<th>Name/type</th>
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<tbody>
<tr>
<td>NSAIDs</td>
<td>Any</td>
</tr>
<tr>
<td>DMARDs</td>
<td>Leflunomide, methotrexate, sulfasalazine, triple non–biologic DMARD (methotrexate, sulfasalazine, hydroxychloroquine)</td>
</tr>
<tr>
<td>Biologics</td>
<td>TNFi</td>
</tr>
<tr>
<td></td>
<td>Adalimumab, etanercept, infliximab, golimumab†</td>
</tr>
<tr>
<td></td>
<td>Non-TNFi‡</td>
</tr>
<tr>
<td></td>
<td>Abatacept (CTLA-4Ig), tocilizumab (anti–interleukin-6 receptor), rituximab (anti–CD20)</td>
</tr>
<tr>
<td>Glucocorticoids</td>
<td>Oral</td>
</tr>
<tr>
<td></td>
<td>Any</td>
</tr>
<tr>
<td></td>
<td>Intraarticular</td>
</tr>
<tr>
<td></td>
<td>Triamcinolone acetonide, triamcinolone hexacetonide, methylprednisolone acetate</td>
</tr>
<tr>
<td>Other interventions</td>
<td>Physical therapy</td>
</tr>
<tr>
<td></td>
<td>Occupational therapy</td>
</tr>
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</table>

* NSAIDs = nonsteroidal antiinflammatory drugs; DMARDs = disease-modifying antirheumatic drugs; TNFi = tumor necrosis factor inhibitor.
† Certolizumab was not included in the recommendations because no pediatric data were yet available.
‡ Evaluated for polyarthritis only.

Literature searches, data abstraction, and rating the quality of evidence. Systematic searches of the published English-language literature included Ovid Medline, PubMed, Embase, and the Cochrane Library (including Cochrane Database of Systematic Reviews, Database of Abstracts of Reviews of Effects, Cochrane Central Register of Controlled Trials, and Health Technology Assessments) from the beginning of each database through June 12, 2017 (see Supplementary Appendix 4, available on the Arthritis Care & Research web site at http://onlinelibrary.wiley.com/doi/10.1002/acr.23870/abstract); updated searches were conducted on October 13, 2017. DistillerSR software (https://distillersr.com/products/distillersr-systematic-reviewsoftware/) facilitated duplicate screening of literature search results. Supplementary Appendix 5, available on the Arthritis Care & Research web site at http://onlinelibrary.wiley.com/doi/10.1002/acr.23870/abstract, shows the citation flow chart. Reviewers entered extracted data into RevMan software (http://tech.cochrane.org/revman) and evaluated the risk of bias of primary studies using the Cochrane risk of bias tool (http://handbook.cochrane.org/). RevMan files were exported into GRADEpro software to formulate a table showing the GRADE summary of findings (Supplementary Appendix 6, available on the Arthritis Care & Research web site at http://onlinelibrary.wiley.com/doi/10.1002/acr.23870/abstract) for each PICO question (31).

When available, the evidence summaries included the benefits and harms for outcomes of interest across studies, the relative effect (95% confidence interval), the number of participants, and the absolute effects. GRADE criteria provided the framework for judging the overall quality of evidence (16). The Literature Review Team rated the quality of evidence for each critical and important outcome as high, moderate, low, or very low quality, taking into account limitations of study design, risk of bias, inconsistency, indirectness, imprecision, and other considerations. The overall quality listed in this report for a body of evidence (i.e., either an individual article or a group of articles) is not a statement about the methodologic quality of the study (or studies). Rather, the intention was to rate the article(s) in relation to the PICO question under consideration. As a result, a very well-conducted study might be rated lower in the evidence report. For example, if the population or intervention being studied does not completely match the population or intervention being examined by the PICO question, the evidence is downgraded for indirectness. The overall quality of evidence may also be downgraded due to imprecision in the effect estimate (e.g., wide confidence intervals or a low number of patients or events).

During the Voting Panel meeting, the panel also considered relevant data from adult studies, but these studies were not systematically searched or compiled in the evidence report. The Voting Panel was provided with a copy of the evidence report from the ACR/Spondylitis Association of America/Spondyloarthritis Research and Treatment Network 2015 Recommendations for the Treatment of Ankylosing Spondylitis and Nonradiographic Axial Spondyloarthritis as a reference (32) (https://www.rheumatology.org/Portals/0/Files/Axial-SpA-Guideline-Supplement-E.pdf).

Moving from evidence to recommendations. Each recommendation was made based on a consideration of the balance of relative benefits and harms of the treatment options under consideration, the quality of the evidence (e.g., confidence in the effect estimates), and patients’ values and preferences, as per GRADE methodology. Discussion points and voting results from the Parent and Patient Panel meeting were presented during the Voting Panel meeting, as relevant. When the literature did not clearly guide recommendations, recommendations were based on the experience of the Voting Panel members (including physicians and the 2 patients present) as well as the results from the Parent and Patient Panel. Financial costs were not formally considered during the voting process.

Consensus building. The Voting Panel voted on the direction and strength of the recommendation related to each PICO question. Recommendations required a 70% level of
agreement; if 70% agreement was not achieved during an initial vote, the panel members held additional discussions before re-voting, including rewording of recommendations if needed, until consensus was attained (33). Discussion and iterative voting occurred until consensus was achieved. An additional round of voting was conducted online after the Voting Panel meeting to address questions that arose during preparation of the final recommendations. For each recommendation, a written explanation is provided, describing the reasons for this decision and conditions under which the alternative choice may be preferable, when relevant.

Moving from recommendations to practice. These recommendations are designed to help health care providers, caregivers, and patients engage in shared decision-making regarding treatment choices. Health care providers, caregivers, and patients must take into consideration not only clinical phenotype and level of disease activity but also comorbidities, response to and tolerance of prior therapies, patient’s values and preferences, and patient’s functional status and functional goals when choosing the optimal therapy for an individual patient at the given point in treatment.

RESULTS/RECOMMENDATIONS

How to interpret the recommendations (18–20)

1. A strong recommendation means that the Voting Panel was confident that the desirable effects of following the recommendation outweigh the undesirable effects (or vice versa), so the course of action would apply to all or almost all patients, and only a small proportion would not want to follow the recommendation. In some cases, strong recommendations were made even in the absence of moderate- or high-quality evidence based on Voting Panel experience and data from adult studies.

2. A conditional recommendation means that the Voting Panel believed that the desirable effects of following the recommendation probably outweigh the undesirable effects, so the course of action would apply to the majority of the patients, but some may not want to follow the recommendation. Because of this, conditional recommendations are particularly preference-sensitive and warrant a shared decision-making approach. Conditional recommendations were generally based on low- to very low-quality evidence. Most recommendations in this guideline are conditional.

3. For each recommendation, Supplementary Appendix 6 (available on the Arthritis Care & Research web site at http://onlinelibrary.wiley.com/doi/10.1002/acr.23870/abstract) provides details regarding the PICO questions and the GRADE evidence tables. PICO questions were combined when possible to create simplified recommendations.

General recommendations for patients with JIA and polyarthritis (Table 3)

For this population, an initial set of general recommendations was made regarding NSAID, DMARD, intraarticular glucocorticoid, and biologic use. These general recommendations are intended to apply to the subsequent specific polyarthritis recommendations addressing these medications, because the recommendations were not anticipated to differ based on initial versus subsequent therapy, level of disease activity, or presence or absence of risk factors. For example, in PICO A.2 and A.3, methotrexate is conditionally recommended over leflunomide and sulfasalazine. It is intended that this recommendation apply to subsequent recommendations referring to DMARD therapy.

Each recommendation in this section is prefaced with the statement “In children and adolescents with JIA and active polyarthritis...”

PICO A.1. NSAIDs are conditionally recommended as adjunct therapy. This recommendation is conditional based on the very low quality of evidence and incorporation of patient and caregiver preferences, particularly concerns regarding medication adverse effects. In general, NSAIDs were thought to be appropriate for symptom management, particularly during initiation or escalation of therapy with DMARDs or biologics (34–36). It was acknowledged that NSAIDs are not appropriate as monotherapy for chronic, persistent synovitis.

PICO A.2–A.3. Using methotrexate is conditionally recommended over leflunomide or sulfasalazine.

Leflunomide. The quality of supporting evidence for this recommendation was moderate. The recommendation to favor methotrexate over leflunomide is due to the greater volume of data supporting the effectiveness of methotrexate. The Voting Panel also specifically noted the lack of data for the dosing, safety, and effectiveness of leflunomide in children younger than age 3 years and lack of long-term safety data for leflunomide in general for children with polyarthritis (37,38). In addition, there currently is no liquid form of this medication, which may make administration difficult, particularly in younger children.

Sulfasalazine. The recommendation for methotrexate over sulfasalazine is conditional, because the supporting evidence is of very low quality, there are no head-to-head comparison studies, and there are more data supporting the effectiveness of methotrexate. There were also concerns raised by the Voting Panel regarding the safety of sulfasalazine as compared to methotrexate, specifically the risk of Stevens-Johnson syndrome and bone marrow suppression (35,36).

Although methotrexate is conditionally recommended over each of these therapies, it is important to note that the Parent and Patient Voting Panel participants stated that they would want to be made aware of available alternatives to methotrexate, because
the adverse effects of methotrexate, particularly gastrointestinal intolerance, are very limiting for some children.

**PICO A.4.** Using subcutaneous methotrexate is conditionally recommended over oral methotrexate. This recommendation is conditional, because the supporting evidence is of very low quality, and patient preferences may guide the choice of route of administration (39–46). The strength of the recommendation also reflects Voting Panel experience, lack of certainty regarding differences in adverse event rates between the 2 routes of administration, consideration of data suggesting variable bioavailability of oral methotrexate (particularly at higher doses), and the goal of optimizing methotrexate effectiveness prior to escalating therapy (47,48).

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**Table 3.** General medication recommendations for children and adolescents with JIA and polyarthritis

<table>
<thead>
<tr>
<th>Recommendation</th>
<th>Level of evidence</th>
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<tbody>
<tr>
<td><strong>NSAIDs</strong></td>
<td></td>
</tr>
<tr>
<td>NSAIDs are conditionally recommended as adjunct therapy (PICO A.1).</td>
<td>Very low</td>
</tr>
<tr>
<td><strong>DMARDs</strong></td>
<td></td>
</tr>
<tr>
<td>Using methotrexate is conditionally recommended over leflunomide or sulfasalazine (PICO A.2, A.3).</td>
<td>Moderate (leflunomide); very low (sulfasalazine)</td>
</tr>
<tr>
<td>Using subcutaneous methotrexate is conditionally recommended over oral methotrexate (PICO A.4).</td>
<td>Very low</td>
</tr>
<tr>
<td><strong>Glucocorticoids</strong></td>
<td></td>
</tr>
<tr>
<td>Intraarticular glucocorticoids are conditionally recommended as adjunct therapy (PICO A.5).</td>
<td>Very low</td>
</tr>
<tr>
<td>Triamcinolone hexacetonide is strongly recommended over triamcinolone acetonide for intraarticular glucocorticoid injections (PICO A.6).</td>
<td>Moderate</td>
</tr>
<tr>
<td>Bridging therapy with a limited course of oral glucocorticoids (&lt;3 months) during initiation or escalation of therapy in patients with high or moderate disease activity is conditionally recommended (PICO A.7).†</td>
<td>Very low</td>
</tr>
<tr>
<td>Bridging therapy may be of most utility in the setting of limited mobility and/or significant symptoms.</td>
<td></td>
</tr>
<tr>
<td>Conditionally recommend against bridging therapy with a limited course of oral glucocorticoids (&lt;3 months) in patients with low disease activity (PICO A.8).</td>
<td>Very low</td>
</tr>
<tr>
<td>Strongly recommend against adding chronic low-dose glucocorticoid, irrespective of risk factors or disease activity (PICO A.9).</td>
<td>Very low</td>
</tr>
<tr>
<td><strong>Biologic DMARDs</strong></td>
<td></td>
</tr>
<tr>
<td>In children and adolescents with JIA and polyarthritis starting treatment with a biologic (etanercept, adalimumab, golimumab, abatacept, or tocilizumab) combination therapy with a DMARD is conditionally recommended over biologic monotherapy (PICO A.10, A.11, A.12, A.13, A.14).</td>
<td>Very low (etanercept, golimumab); low (adalimumab); moderate (abatacept, tocilizumab)</td>
</tr>
<tr>
<td>Combination therapy with a DMARD is strongly recommended for infliximab (PICO A.15).</td>
<td>Low</td>
</tr>
<tr>
<td><strong>Physical therapy and occupational therapy</strong></td>
<td></td>
</tr>
<tr>
<td>In children and adolescents with JIA and polyarthritis who have or are at risk of functional limitations, using physical therapy and/or occupational therapy is conditionally recommended (PICO A.16, PICO A.17).</td>
<td>Low (physical therapy); very low (occupational therapy)</td>
</tr>
</tbody>
</table>

* JIA = juvenile idiopathic arthritis; NSAIDs = nonsteroidal antiinflammatory drugs; PICO = Patient/Population, Intervention, Comparison, and Outcomes; DMARDs = disease-modifying antirheumatic drugs.
† A bridging course of oral glucocorticoids was defined as a short course (<3 months) of oral glucocorticoids intended to control disease activity quickly during the initiation or escalation of therapy. An adequate trial of methotrexate was considered to be 3 months. If no or minimal response is observed after 6–8 weeks, it was agreed that changing or adding therapy may be appropriate.
PICO A.6. Triamcinolone hexacetonide is strongly recommended over triamcinolone acetonide for intraarticular glucocorticoid injections. The quality of supporting evidence for this recommendation was moderate (50). This recommendation was further supported by observational studies showing improved outcomes with triamcinolone hexacetonide in oligoarticular JIA and rheumatoid arthritis (RA) (51,52). Voting panel members specifically noted their consistent and repeated observation of more complete and longer duration of clinical response without increased adverse effects with triamcinolone hexacetonide versus triamcinolone acetonide. The Parent and Patient Voting Panel also supported this judgment on the strength of the recommendation.

PICO A.7. Bridging therapy with a limited course of oral glucocorticoids (<3 months) during initiation or escalation of therapy in patients with high or moderate disease activity is conditionally recommended. This recommendation is conditional based on very low quality of supporting evidence and the known risks associated with systemic glucocorticoid treatment. Parents and patients agreed that bridging therapy was acceptable in this setting. Bridging therapy with glucocorticoids may have most utility
in the setting of high disease activity, limited mobility, and/or significant symptoms.

PICO A.8. Conditionally recommend against bridging therapy with a limited course of oral glucocorticoids (<3 months) in patients with low disease activity. The quality of evidence for this recommendation was very low. Intraarticular glucocorticoid injection was considered preferable in this setting.

PICO A.9. Strongly recommend against adding chronic low-dose glucocorticoids, irrespective of risk factors or disease activity. The quality of supporting evidence for this recommendation was very low. The recommendation was strong, however, given the known adverse effects of long-term systemic glucocorticoid treatment in children, particularly growth suppression, weight gain, osteopenia and cataracts, and availability of other treatment options. The Voting Panel agreed that in the setting of low disease activity, targeted joint injections may be more appropriate (PICO A.5). In the setting of moderate or high disease activity, escalating DMARD or biologic therapy is likely more appropriate.

PICO A.10–A.14. In children and adolescents with JIA and polyarthritis initiating treatment with a biologic (etanercept, adalimumab, golimumab, abatacept, or tocilizumab), combination therapy with a DMARD is conditionally recommended over biologic monotherapy. This recommendation is intended for patients initiating biologics for additional disease control and not intended for patients who may be tapering therapy due to inactive disease, for whom tapering or removal of the DMARD while continuing biologic therapy may be an appropriate strategy. The available evidence addresses combination therapy with methotrexate, with no evidence identified for other DMARDs. There was variability in the quality of supporting evidence for the medications, ranging from very low (etanercept, golimumab) to low (abatacept or tocilizumab) and moderate (adalimumab) (39,53–67). The potential benefit of methotrexate treatment for prevention of antidrug antibodies to adalimumab was included in the discussion for concomitant DMARD use with that medication (61). The overall recommendation is conditional based on the quality of supporting evidence and variable parent and patient preferences given the burden of taking multiple medications and concerns regarding methotrexate intolerance. The Voting Panel recognized that there may be situations in which biologic monotherapy is also acceptable, particularly in the setting of adequate disease control or methotrexate intolerance.

PICO A.15. Combination therapy with a DMARD is strongly recommended for infliximab. Using infliximab in combination with a DMARD was a strong recommendation despite the low quality of evidence, primarily given more extensive experience with the need for combination therapy to reduce the risk of antidrug antibody formation (68,69).

PICO A.16, A.17. In children and adolescents with JIA and polyarthritis who have or are at risk for functional limitations, using physical therapy (PT) and/or occupational therapy (OT) is conditionally recommended. This recommendation is conditional based on the low quality of supporting evidence for PT, very low level of evidence for OT, and Voting Panel experience (70,71).

Recommendations for the initial and subsequent treatment of JIA and polyarthritis (Table 4 and Figure 1)

While the initial set of PICO questions for polyarthritis also included comparisons between specific biologics, these questions were discarded at the in-person Voting Panel meeting due to lack of evidence to guide decision-making in those scenarios. Although there is most experience with tumor necrosis factor inhibitors (TNFi) as the initial biologic, the class of initial biologic is not specified in the recommendations, again due to lack of comparative data, and the consideration that non-TNFi biologics may be preferred in certain scenarios based on patient-level factors (e.g., family history of demyelinating disease) and preferences. In the recommendations (see below), biologic therapy refers to TNFi, abatacept, or tocilizumab, with the exception of PICO B.9, in which rituximab is also addressed. Each recommendation in this section is prefaced with the statement “In children and adolescents with JIA and active polyarthritis…”

Initial therapy

PICO B.1 Initial therapy with a DMARD is strongly recommended over NSAID monotherapy. This recommendation is strong based on moderate quality of supporting evidence, known risk of permanent joint damage associated with ongoing, active disease, and Voting Panel experience (35,36,41).

PICO B.2. Using methotrexate monotherapy as initial therapy is conditionally recommended over triple DMARD therapy. The quality of evidence for this recommendation was low, based on the available trial being relatively small and not blinded (72). Parents and patients also expressed concerns about the burden of taking the 3 different medications, but they did state a preference to be informed about this treatment option.

PICO B.3. For patients without risk factors, initial therapy with a DMARD is conditionally recommended over a biologic. This recommendation is conditional based on low quality of supporting evidence and parent and patient differing preferences regarding the risks and benefits of DMARDs and biologics. While initial treatment with a biologic has been studied in TREAT-JIA (Trial of Early Aggressive Therapy in Polyarticular JIA) and ACUTE-JIA (Aggressive Combination Drug Therapy in Early Polyarticular JIA), the results were not thought to be conclusive enough to support
There are situations where initial therapy that includes a DMARD is conditionally recommended over a biologic, recognizing that there are situations where initial therapy that includes biologics as initial therapy for low-risk patients (40,72). Of note, the majority of the Parent and Patient Panel voted against DMARDs as initial therapy, because a number of the participants had considerable adverse effects with methotrexate and had experienced better outcomes with biologics.

**PICO B.4.** For patients with risk factors, initial therapy with a DMARD is conditionally recommended over a biologic, recognizing that there are situations where initial therapy that includes a biologic may be preferred. This recommendation is conditional based on the low quality of supporting evidence and parent and patient differing preferences regarding the risks and benefits of DMARDs and biologics. While initial treatment with a biologic has been studied in TREAT-JIA and ACUTE-JIA, the results were not thought to be conclusive enough to support biologics as initial therapy (40,72). However, the Voting Panel acknowledged that biologics may be appropriate initial therapy for some patients with risk factors and involvement of high-risk joints (e.g.,

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<table>
<thead>
<tr>
<th>PICO B.4.</th>
<th>Recommendations</th>
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| **For patients with risk factors, initial therapy with a DMARD is conditionally recommended over a biologic, recognizing that there are situations where initial therapy that includes a biologic may be preferred.** This recommendation is conditional based on the low quality of supporting evidence and parent and patient differing preferences regarding the risks and benefits of DMARDs and biologics. While initial treatment with a biologic has been studied in TREAT-JIA and ACUTE-JIA, the results were not thought to be conclusive enough to support biologics as initial therapy (40,72). However, the Voting Panel acknowledged that biologics may be appropriate initial therapy for some patients with risk factors and involvement of high-risk joints (e.g.,

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**Figure 1.** Summary of primary recommendations for the initial and subsequent treatment of children with juvenile idiopathic arthritis (JIA) and active polyarthritis (see also Tables 3 and 4; for patients with sacroiliitis and/or enthesitis, see also Tables 5 and 6). The clinical Juvenile Arthritis Disease Activity Score based on 10 joints (cJADAS-10) was used to define low disease activity (≤2.5 with ≥1 active joint) versus moderate/high disease activity (>2.5). Although it is provided as a general parameter, the cJADAS-10 should be interpreted within the clinical context. An adequate trial of methotrexate was considered to be 3 months. If no or minimal response is observed after 6–8 weeks, it was agreed that changing or escalating therapy may be appropriate. Shared decision-making between the physician, parents, and patient, including discussion of recommended treatments and potential alternatives, is recommended when initiating or escalating treatment. The Patient/Population, Intervention, Comparison, and Outcomes (PICO) questions are shown in brackets, and quality of evidence is shown in parentheses. DMARD = disease-modifying antirheumatic drug; NSAID = nonsteroidal antiinflammatory drug; PT = physical therapy; OT = occupational therapy; TNFi = tumor necrosis factor inhibitor.

<table>
<thead>
<tr>
<th><strong>Low disease activity, biologic naive</strong></th>
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<tbody>
<tr>
<td><strong>Initial therapy:</strong> Methotrexate</td>
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<tr>
<td>- Subcutaneous over oral methotrexate [A.4] (very low)</td>
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<tr>
<td>- Methotrexate over sulfasalazine [A.3] (very low)</td>
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<tr>
<td>- Methotrexate over leflunomide [A.2] (moderate)</td>
</tr>
<tr>
<td>- Single DMARD over triple DMARD therapy [B.2] (low)</td>
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<tr>
<td>- DMARD over biologic [B.3-4] (low)*</td>
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**Legend**

- Strongly recommend
- Conditionally recommend
- Conditionally recommend against
- Strongly recommend against
cervical spine, hip, and wrist), high disease activity, and/or for those judged by their physician to be at high risk of disabling joint damage. Of note, the majority of the Parent and Patient group voted against DMARDs as initial therapy, because a number of the participants had considerable adverse effects with methotrexate and had experienced better outcomes with biologics. Despite a DMARD or biologic, escalating therapy is conditionally recommended over no escalation of therapy. This recommendation is conditional based on the very low quality of supporting evidence and parent and patient preferences regarding the risks and benefits of the treatment options. For this recommendation, escalating therapy is defined as any of the following: intraarticular glucocorticoid injection, increasing the DMARD or biologic dose (if not at optimal dosage), or changing biologic. Changing to an alternate DMARD (methotrexate) was suggested primarily for patients who had not yet received methotrexate and had not yet escalated to treatment with a biologic. Additional considerations included a degree of improvement while receiving current therapy.

**Subsequent therapy in patients with low disease activity (cJADAS-10 ≤2.5 and at least 1 active joint)**

*PICO B.5, B.6. In patients with JIA and polyarthritis and low disease activity (cJADAS-10 ≤2.5 and at least 1 active joint)*

<table>
<thead>
<tr>
<th>Table 5. Recommendations for the initial and subsequent treatment of children and adolescents with JIA and sacroiliitis*</th>
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<tbody>
<tr>
<td><strong>Recommendation</strong></td>
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<tr>
<td>In children and adolescents with active sacroiliitis, treatment with an NSAID is <em>strongly</em> recommended over no treatment with an NSAID (PICO C.1).</td>
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<tr>
<td>In children and adolescents with active sacroiliitis despite treatment with NSAIDs:</td>
</tr>
<tr>
<td>• Adding TNFi is <em>strongly</em> recommended over continued NSAID monotherapy (PICO C.2).</td>
</tr>
<tr>
<td>• Using sulfasalazine for patients who have contraindications to TNFi or have failed more than one TNFi is conditionally recommended (PICO C.3).</td>
</tr>
<tr>
<td>• <em>Strongly</em> recommend against using methotrexate monotherapy (PICO C.4).</td>
</tr>
</tbody>
</table>

**Glucocorticoids**

In children and adolescents with active sacroiliitis despite treatment with NSAIDs:

• Bridging therapy with a limited course of oral glucocorticoids (<3 months) during initiation or escalation of therapy is conditionally recommended (PICO C.5).
  Bridging therapy may be of most utility in the setting of high disease activity, limited mobility, and/or significant symptoms.
• Intraarticular glucocorticoid injection of the sacroiliac joints as adjunct therapy is conditionally recommended (PICO C.6).

**Physical therapy**

• In children and adolescents with sacroilitis who have or are at risk for functional limitations, using physical therapy is conditionally recommended (PICO C.7).

* TNFi = tumor necrosis factor inhibitor (etanercept, adalimumab, infliximab, golimumab) (see Table 3 for other definitions).
† A bridging course of oral glucocorticoids was defined as a short course (<3 months) of oral glucocorticoids intended to control disease activity quickly during the initiation or escalation of therapy.

**Table 6. Recommendations for the initial and subsequent treatment of children and adolescents with JIA and enthesitis* |

| **Recommendation** | **Level of evidence** |
|-------------------------------------------------|
| In children and adolescents with active enthesitis, NSAID treatment is *strongly* recommended over no treatment with an NSAID (PICO D.1). | Very low |
| In children and adolescents with active enthesitis despite treatment with NSAIDs: | |
| • Using a TNFi is conditionally recommended over methotrexate or sulfasalazine (PICO D.2, D.3). | Low |
| • Bridging therapy with a limited course of oral glucocorticoids (<3 months) during initiation or escalation of therapy is conditionally recommended (PICO D.4).† | Very low |
  Bridging therapy may be of most utility in the setting of high disease activity, limited mobility, and/or significant symptoms.
| **Physical therapy** | |
| • In children and adolescents with enthesitis who have or are at risk for functional limitations, using physical therapy is conditionally recommended (PICO D.5). | Very low |

* TNFi = tumor necrosis factor inhibitor (etanercept, adalimumab, infliximab, golimumab) (see Table 3 for other definitions).
† A bridging course of oral glucocorticoids was defined as a short course (<3 months) of oral glucocorticoids intended to control disease activity quickly during the initiation or escalation of therapy.
and the specific joint that was active. Synovitis preventing ambulation or otherwise interfering with important daily activities was identified as a factor that would guide more aggressive intervention (e.g., intraarticular glucocorticoid injection or adding/changing biologic).

**Subsequent therapy in patients with moderate or high disease activity (cJADAS-10 >2.5)**

**PICO B.7. In patients with JIA and polyarthritis and moderate or high disease activity despite DMARD monotherapy, adding a biologic to the original DMARD is conditionally recommended over changing to a second DMARD.** This recommendation was conditional based on the low quality of supporting evidence and parent and patient preferences regarding the risks and benefits of DMARDs and biologic medications (29,58,61,64,65,67,72–82).

**PICO B.8. In patients with JIA and polyarthritis and moderate or high disease activity receiving DMARD monotherapy, adding a biologic is conditionally recommended over changing to triple DMARD therapy.** The quality of evidence for this recommendation was low, based on the published pediatric trial being relatively small and not blinded (72). Although studies in RA have suggested that triple therapy is non-inferior to biologic therapy, the pediatric study showed improved outcomes for biologic therapy over triple DMARD therapy (83,84). This recommendation was also supported by Voting Panel concerns about adherence to the regimen and tolerability and parent and patient concerns about the burden of taking the 3 different medications.

**PICO B.9. In patients with JIA and polyarthritis and moderate or high disease activity receiving a first TNFi (with or without DMARD), switching to a non-TNFi biologic (tocilizumab or abatacept) is conditionally recommended over switching to a second TNFi.** This recommendation was conditional based on very low quality of supporting evidence and parent and patient preferences regarding the risks and benefits of biologics with different mechanisms of action (79,85). In making this recommendation, the Voting Panel also considered data from RA that have suggested better outcomes with switching to a non-TNFi biologic (86–88). The Voting Panel agreed that a second TNFi may be appropriate for patients who had a good initial response to their first TNFi (i.e., secondary failure), particularly failure due to the presence of suspected or measured antidrug antibodies to the first TNFi.

**PICO B.10. In patients with JIA and polyarthritis and moderate or high disease activity despite a second biologic, using a TNFi, abatacept, or tocilizumab (depending upon prior biologics received) is conditionally recommended over rituximab.** This recommendation was conditional based on very low quality of supporting evidence, Voting Panel member experience, and parent and patient preferences regarding the risks and benefits of biologics with different mechanisms of action. This recommendation was also supported by the availability of data from randomized clinical trials of tocilizumab and abatacept establishing their efficacy in JIA, which is lacking for rituximab (64,65,67). In addition, the article identified for the evidence report showed a higher rate of serious adverse events for rituximab compared to other biologics (78). The Voting Panel did discuss that rituximab may be considered earlier for RF-positive children based on data from RA, although the other 3 classes of biologics would still be primarily recommended (89).

**Recommendations for the treatment of JIA and sacroiliitis (Table 5)**

**PICO C.1. In children and adolescents with JIA and active sacroiliitis, treatment with an NSAID is strongly recommended over no treatment with an NSAID.** This recommendation was strong despite very low quality of supporting evidence in children, given the established utility of NSAIDs in adult spondyloarthritis and the analgesic effects of NSAIDs in children with other forms of arthritis. This recommendation is in alignment with the treatment recommendations for ankylosing spondylitis co-developed by the American College of Rheumatology/Spondylitis Association of America Spondyloarthritis Research and Treatment Network (32).

**PICO C.2. In children and adolescents with active sacroiliitis despite NSAIDs, adding a TNFi is strongly recommended over continued NSAID monotherapy.** Although the quality of supporting evidence in pediatrics for this recommendation was low, this recommendation is based on evaluation of both pediatric data and data from adult spondyloarthritis that include randomized controlled trials showing benefit (80,90–99).

**PICO C.3. In children and adolescents with active sacroiliitis despite NSAIDs, using sulfasalazine for patients who have contraindications to TNFi or have failed more than one TNFi is conditionally recommended.** This recommendation was conditional based on low quality of supporting evidence, particularly the relatively limited efficacy of sulfasalazine demonstrated in a randomized controlled trial in juvenile spondyloarthritis (100). However, it was considered as a potential option for patients with contraindications to TNFi and based on parent and patient considerations of the risks and benefits of biologics versus sulfasalazine. Sulfasalazine was also considered as an option for patients with adverse events associated with their initial TNFi that would be considered a class effect and who would therefore not be able to receive additional TNFi. Non-TNFi biologics...
(e.g., interleukin-17 [IL-17] inhibitors) were not considered by the Voting Panel, because there are no published pediatric studies.

**PICO C.4.** In children and adolescents with active sacroiliitis despite treatment with NSAIDs, strongly recommend against using methotrexate monotherapy. Although the quality of supporting evidence for this recommendation was very low, this recommendation is based on data from adult spondyloarthritis suggesting lack of effectiveness (92,101–103). While we recommend against using methotrexate as a treatment for sacroiliitis, methotrexate may have utility as adjunct therapy in patients with concomitant peripheral polyarthritis or to prevent the development of antidrug antibodies against monoclonal TNFis.

**PICO C.5.** In children and adolescents with active sacroiliitis despite treatment with NSAIDs, bridging therapy with a limited course of oral glucocorticoids (<3 months) during initiation or escalation of therapy is conditionally recommended. This recommendation is conditional based on very low quality of supporting evidence and known risks of glucocorticoid treatment. Bridging therapy with oral glucocorticoids may have the most utility in the setting of high disease activity, limited mobility, and/or significant symptoms.

**PICO C.6.** In children and adolescents with active sacroiliitis despite treatment with NSAIDs, intraarticular glucocorticoid injection of the sacroiliac joints as adjunct therapy is conditionally recommended. This recommendation was conditional based on very low quality of evidence and on varying parent and patient preferences regarding the procedure.

**PICO C.7.** In children and adolescents with sacroiliitis who have or are at risk for functional limitations, using PT is conditionally recommended. This recommendation was conditional based on the very low quality of supporting evidence and Voting Panel experience. It was also discussed that there may be a role for PT and activity modification in specifically identifying and reducing mechanical factors contributing to microtrauma and repetitive stress that could potentially contribute to disease activity in these patients (104).

**Recommendations for the treatment of JIA and enthesitis (Table 6)**

**PICO D.1.** In children and adolescents with JIA and active enthesitis, NSAID treatment is strongly recommended over no treatment with an NSAID. This recommendation is strong despite the very low quality of supporting evidence based on Voting Panel experience, established analgesic effects, and data from adult disease showing benefit (105).

**PICO D.2, D.3.** In children and adolescents with JIA and active enthesitis despite treatment with NSAIDs, using a TNFi is conditionally recommended over methotrexate or sulfasalazine. This recommendation is conditional based on the low quality of supporting evidence. While TNFi is preferred, the Voting Panel discussed that a trial of methotrexate or sulfasalazine may be warranted for patients with contraindications to TNFi, patients with mild enthesitis, and patients with concomitant active peripheral polyarthritis (80,90–95,100).

**PICO D.4.** In children and adolescents with JIA and chronic active enthesitis despite treatment with NSAIDs, bridging therapy with a limited course of oral glucocorticoids (<3 months) during initiation or escalation of therapy is conditionally recommended. This recommendation is conditional based on very low quality of supporting evidence and known risks of glucocorticoid treatment in the pediatric population. Bridging therapy with glucocorticoids may have most utility in the setting of high disease activity, limited mobility, and/or significant symptoms.

**PICO D.5.** In children and adolescents with JIA and enthesitis who have or are at risk for functional limitations, using PT is conditionally recommended. This recommendation was conditional based on very low quality of evidence and Voting Panel experience.

**DISCUSSION**

This guideline includes 39 recommendations for the treatment of children with JIA and non-systemic polyarthritis, sacroiliitis, and enthesitis. The quality of most of the available evidence was low or very low in relation to the relevant clinical PICO questions, resulting in 31 of the recommendations being conditional.

These recommendations provide an updated approach to the treatment of children with non-systemic polyarthritis, sacroiliitis, and enthesitis. These populations were chosen for this guideline because they have been the focus of significant recent research, with better delineation of the underlying biology and additional treatments available since the 2011 ACR recommendations for JIA. Similar to the 2011 recommendations, this guideline defined patient populations by clinical phenotypes rather than ILAR categories. This decision was made because data continue to suggest that current JIA categories may not accurately reflect the underlying biology and anticipated treatment responses in patients with juvenile arthritis.

This guideline differs from the 2011 recommendations in the definitions of risk factors and disease activity assessment used to generate patient scenarios. Although PICO questions were initially stratified by risk factors and disease activity, the Voting Panel ultimately determined that in most scenarios there were not sufficient data to recommend different treatments for these
patients, and recommendations were frequently combined. Another important difference from the 2011 recommendations is that initial NSAID monotherapy for polyarthritis is no longer recommended, given the established benefits of early initiation of DMARD treatment. Individual PICO questions for each biologic were initially considered but subsequently dropped by the Voting Panel, given mostly equivalent data for safety and efficacy between the biologics and lack of head-to-head comparisons. The exceptions were that TNFi are specifically recommended for sacroiliitis, and rituximab is considered only after TNFi, abatacept, and tocilizumab have been tried. This approach has resulted in a simplified treatment algorithm. Last, this guideline also includes recommendations for escalating care in the setting of low disease activity, highlighting the importance of achieving and maintaining complete disease control, which was not previously addressed.

The current recommendations also differ from the 2011 recommendations in that they were developed using GRADE methodology instead of the RAND/UCLA Appropriateness Method. The systematic, transparent, and explicit process of developing recommendations through GRADE is a major feature accelerating its adoption by professional groups internationally (www.gradeworkinggroup.org). Important features of this method are 1) specification of the patient groups, interventions, competing alternatives, and outcomes so that each recommendation is clearly focused on a particular clinical situation; 2) grading of the quality of evidence; and 3) basing the strength of recommendations on the quality of evidence, balance of benefits and harms, and patient values and preferences for different treatment options. This guideline considered parent and patient preferences assessed by a separate Parent and Patient Panel. Primary themes that emerged from that discussion were: 1) the importance of shared decision-making; 2) the importance of parents and patients receiving information regarding not only the preferred medication or intervention but also the alternatives; and 3) parent/patient support of earlier consideration of biologics given their experiences with decreased adverse effects and improved quality of life with the use of these medications relative to their experiences with methotrexate. Although this was a select group of parents and patients, and their experiences may not be representative of all patients, their discussion provided an important perspective that was incorporated into the Voting Panel discussion and voting.

A topic of particular debate among the Voting Panel was the appropriateness of the use of biologics as initial therapy in children with polyarthritis, particularly for those with risk factors. Ultimately, non–biologic DMARD therapy was recommended, but it was noted that there may be some patients for whom initial biologic therapy is indicated. This remains an area of active research, and currently ongoing studies may better clarify which patients are most likely to benefit from initial biologic therapy. Importantly, studies in pediatrics are underway or planned for a number of new medications, including JAK inhibitors and IL-17 and IL-12/23 inhibitors, and these medications may become useful additions as treatment options for JIA, particularly in patients with sacroiliitis for whom limited options exist. Other studies currently underway in parallel adult diseases may also inform the optimal treatment of enthesitis and the treatment of peripheral spondyloarthritis. Future guideline efforts may determine where these treatments fit into the treatment algorithm and will incorporate the results from the ongoing studies once complete.

Nonpharmacologic interventions addressed in this guideline included PT and OT. In each case, very limited data were identified, and future research in these modalities will be helpful in identifying patients most likely to benefit from these interventions and which modalities have most effectiveness for particular clinical scenarios.

The cJADAS-10 was used to provide general disease activity parameters for defining low and moderate/high disease activity. However, this is not intended to be prescriptive and should be interpreted within the overall clinical context. Furthermore, because the JADAS is a relatively new disease activity measurement tool, new cutoffs may be proposed, since additional data are generated that may accommodate different numbers of active joints or other levels of physician’s or parent’s global assessment. The identification of valid and practical disease activity measures in JIA remains an important research agenda in pediatric rheumatology. Nevertheless, it was agreed by the Voting Panel that treatment should be escalated in patients with even 1 active joint. Formal recommendations regarding disease activity measurement tools, cutoffs, and monitoring intervals were not specifically addressed in this guideline. Last, the management of inactive disease and the tapering and withdrawal of medications for patients with inactive disease are not addressed in this guideline but will be important for future guidelines.

Because the quality of evidence was overall low and most recommendations were conditional, clinicians, caregivers, and patients should use a shared decision-making process when considering these recommendations. While these recommendations are intended to address common clinical situations, all treatment decisions must be individualized, with consideration of the unique aspects of each patient’s presentation, medical history, and preferences.

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AUTHOR CONTRIBUTIONS

All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be published. Dr. Ringold had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Study conception and design. Ringold, Angeles-Han, Beukelman, Lovell, Cuello, Becker, Colbert, Feldman, Ferguson, Gevenater, Guzman, Horonjeff, Ngovic, Schneider, Halaybar, Turner, Reston.

Acquisition of data. Ringold, Angeles-Han, Beukelman, Lovell, Cuello, Becker, Colbert, Feldman, Ferguson, Gevenater, Guzman, Horonjeff, Ngovic, Ombrello, Passo, Rabinovich, Schneider, Halaybar, Hays, Shah, Sullivan, Szymanski, Turungbaev, Reston.

Analysis and interpretation of data. Ringold, Angeles-Han, Beukelman, Lovell, Cuello, Becker, Colbert, Feldman, Ferguson, Gevenater, Guzman, Horonjeff, Ngovic, Ombrello, Passo, Stoll, Schneider, Halaybar, Hays, Shah, Sullivan, Turungbaev, Reston.

REFERENCES


mg/kg in a multicentre 12 week trial in active polyarticular course juvenile idiopathic arthritis. Rheumatology (Oxford) 2009;48:916–9.


