Improving the quality of care in children with juvenile idiopathic arthritis: A step in the right direction

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Editorial

Improving the Quality of Care in Children with Juvenile Idiopathic Arthritis: A Step in the Right Direction

Juvenile idiopathic arthritis (JIA) is heterogeneous in both its clinical presentation and etiology\(^1,2\). JIA can result in permanent physical disability due to joint damage, while additional morbidity can be related to its treatment and to effects on growth and development. Children with inadequately treated or recalcitrant JIA may have chronic pain, mood disturbances, and difficulty with peer relationships, school performance, and attainment of educational and vocational goals\(^3,4,5,6,7,8\). Improved health related quality of life (HRQOL) and physical function are increasingly recognized as key treatment goals because of the influence of this illness on all aspects of a child’s life\(^9\). Accurate evaluation of this multidimensional and heterogeneous disease is challenging, but crucial to improve the quality of care and outcomes in JIA.

Over the last decade there has been an increasing commitment to the assessment and improvement of quality of care following the report of The Institute of Medicine (IOM) Committee on Quality of Healthcare in America, “To Err is Human”\(^10\). The American College of Rheumatology (ACR) and American Board of Pediatrics have charged Pediatric Rheumatology with developing quality measures (QM) for JIA. The initial work of this group included use of an online Delphi survey technique with participation from physicians, advanced practice nurses, and parents of children with JIA for the selection process of QM. In 2008, preliminary QM were published\(^11\). A nominal group technique was used to reach consensus on the proposed set of QM for the process of care. These include 4 broad domains: disease control, safety monitoring, access and relationship (patient/family satisfaction with healthcare)\(^12\). Within several of the QM, the use of validated, reliable, age-appropriate tools for measurement of pain, HRQOL, functional ability, self-efficacy, and parent/patient satisfaction is proposed. A collaborative network, “Pediatric Rheumatology Care and Outcomes Improvement Network (PR-COIN),” has recently been established to develop and evaluate specific disease management strategies to improve the care of children with JIA and to determine how best to incorporate these strategies into clinical practice.

In this issue of The Journal, Filocamo, et al describe the Juvenile Arthritis Multidimensional Assessment Report (JAMAR), an instrument for assessment of children with JIA for use in standard clinical care\(^13\). The authors proposed that this report may help enhance the quality of care of children with JIA by addressing “efficiency” and “efficacy,” two of the quality domains of the “STEEEP” acronym outlined by the IOM (safe, timely, effective, efficient, equitable, patient-centered)\(^14\). One of the novel aspects of this tool is the inclusion of patient-reported outcomes (PRO), in a single tool including morning stiffness, medication side effects, self-report of articular symptoms, and patient satisfaction rating\(^13\). The inclusion of PRO is important to improve quality of care as these are part of the “patient-centered care” domain of quality outlined by the IOM.

Patient-specific indices offer the advantage of identifying salient issues for each patient and are more likely to focus medical attention on the relevant issues; however, they also present unique challenges to ensure the item/measure is reliable, valid, and provides useful information. Without standardization of the items, the response scales do not have the same meaning for each patient. This can make it difficult to understand the numeric meaning of a score. Second, using the data beyond intraindividual comparison is problematic\(^15\). Further studies are needed to improve our understanding of how to interpret the numeric scores of patient-specific indices at both the individual and group level.

The JAMAR contains two measurement tools, the Juvenile Arthritis Functionality Scale (JAFS)\(^16\) and the Pediatric Rheumatology Quality of Life Scale\(^17\), which were developed only recently. For both tools, initial validation studies have been reported. Although research on quality of care in pediatric rheumatology is in its infancy, there has been a longstanding focus on the measurement and reporting of disease outcome, HRQOL, and functional outcome in JIA. There are many tools available to measure clinical response (i.e., ACR Pediatric 30\(^18\)), functional status (i.e., CHAQ\(^19\)), and quality of life (PedsQL\(^20\), Child

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In summary, JIA is a heterogeneous and multidimensional disease. Improving the process of quality care delivery is a topical issue. The development of the JAMAR importantly illustrates the breadth and potential content in a multidimensional report. Future work should be focused on refining and validating the existing measures used in JIA both in clinical practice and in the research setting. Particular attention needs to be focused on interpreting scores/summary scales for PRO. An international consensus on disease activity measures, functional assessment, assessment of HRQOL, and PRO is key to advancing work in this area. In order to evaluate the QM for the process of care, universal agreement on measures to be used will facilitate implementation of QM into routine clinical practice for clinicians. The aim should be to minimize duplication of work and focus on implementation of QM to improve the process of care, then to evaluate these tools and refine them in the plan-do-study-act rapid cycles of improvement.

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