Pediatric rheumatology: addressing the transition to adult-orientated health care

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Abstract: The transition from pediatric to adult health care is often a challenging process due to multiple interwoven complexities, especially for children with chronic medical conditions. Health care transition (HCT) is a process of moving from a pediatric to an adult model of health care with or without a transfer to a new clinician. This paper focuses on what is known about HCT for youth and young adults (Y/YA) with rheumatic diseases within a larger context of HCT recommendations. HCT barriers for youth, families, and providers and current evidence for a structured HCT processes are reviewed. Practical advice is offered on how to approach transition for Y/YA, what tools are available to assist in a successful transition process, and what are the areas of future research that are needed to improve the HCT evidence base.

Keywords: health care transition, pediatric rheumatology, special health care needs, transition, transition readiness assessment

Introduction and background

The transition from pediatric to adult health care is often a challenging process due to multiple interwoven complexities, especially for children with chronic medical conditions. Historically, transition has been characterized by an unsystematic or disorganized transfer of care. Health care transition (HCT) is not just the act of transfer but is the process over time of moving from a pediatric to an adult model of health care with or without transfer to a new clinician. The goals of transition are 1) to improve the ability of youth and young adults (Y/YA) to manage their own health and effectively use health services and 2) to have an organized clinical process in pediatric and adult practices to facilitate transition preparation, transfer of care, and integration into adult-centered care. Patients with special health care needs (SHCN) require increased care and attention, making the transition process more complex. This paper focuses on what is known about HCT for Y/YA with rheumatic diseases within a larger context of HCT recommendations and does not discuss education or vocation, two other important transition components.

It is estimated that there are 18 million adolescents in the USA, of which ~4.5 million have chronic medical conditions. The need for a coordinated systematic transition of care from pediatric to adult providers has increased as advances in health care have facilitated decline in morbidity and mortality of children with chronic medical conditions. More than 90% of children with SHCN are now projected to survive to adulthood and require transitional services.
Both healthy children and those with chronic illnesses experience multiple barriers to successful transition. One such barrier is the lack of transition support that youths experience in transition of care. The 2016 National Survey for Children’s Health of parents of Y/YA aged 12–17 years reports that 83% of youth with SHCN and 86% of youth without SHCN did not receive adequate transition support. Other surveys analyzing transitional needs of youth with juvenile idiopathic arthritis (JIA) and their parents reported a negative transfer experience and stressed the need for increased coordination of care, provider continuity, visit privacy, and an age-appropriate approach to transition of care. A systematic review of patients’ attitudes and experiences of transitional care indicated that some patients feel de-personalized, abandoned, ill-prepared, and not in control of their transition process.

During transition to adult care, there is also a tendency toward increased disease activity. Approximately half of young people with juvenile onset rheumatic arthritis enter adulthood with active disease or develop disease flares as YA. In patients with childhood onset of systemic lupus erythematosus (SLE), there is evidence of increased disease activity in the year following transition. Patients with chronic rheumatic diseases often require long-term treatment with immunosuppressive therapies and are at significant risk of increased disability and morbidity if their care is interrupted during transition. Studies suggest that patients transitioning to an adult rheumatology practice experience significant delays in their first adult visit, which may adversely impact medication adherence and clinic visit show rates. Unfortunately, there is evidence that up to half of young patients do not complete a successful transfer to adult rheumatology, increasing their risk of poor outcomes. In addition to experiencing gaps in care during transfer, youths are at risk of withdrawing from health care services altogether. Furthermore, patients often need increased emotional and mental health support during the transition period. For instance, youths with SLE have shown notable anxiety and depression in the post-transition period. Last, many adolescents and young adults lack important self-management and health care utilization skills needed for the transfer to adult care.

Lack of clinicians’ knowledge regarding transition processes and available resources, decreased access to patient information, and lack of adequate communication, funding, and time are all additional barriers to transition. A survey of pediatric rheumatologists in the Childhood Arthritis and Rheumatology Research Alliance and a survey of transition practices and attitudes among adult rheumatologists in the American College of Rheumatology (ACR) revealed low provider familiarity with transition resources and recommendations, inadequate training in the transition issues of YA, and lack of the necessary time and available personnel to facilitate the transition process. Insufficient communication between pediatric and adult providers can impede successful transition and decrease the quality of patient care. As part of the ACR survey, nearly half of the adult providers reported dissatisfaction with the patient information they received from the pediatric provider. In the USA, funding of transition services proves to be another barrier to successful transfer of care. Last, the implementation of a systematic transition process requires a significant dedication of clinician and care-manager time, especially for youth with socially and medically complex conditions.

**Transition process for practices/systems**

Studies have shown that having a structured transition method can significantly improve transition processes and outcomes. Currently, there are multiple transition models which, although variable in their structure and provider roles, often align across broad categories of processes to improve transition for youth with and without SHCN. These HCT processes are 1) preparation, 2) transfer of care, and 3) integration into adult health care.

In 2011, the American Academy of Pediatrics (AAP), the American Academy of Family Practitioners (AAFP), and the American College of Physicians (ACP) joined together to outline their suggested HCT approach in a clinical report called Supporting the Health Care Transition from Adolescence to Adulthood in the Medical Home. It provides an approach to HCT including the need to assess transition skill readiness and patient education, preparation of a medical summary, and communication between the pediatric and the adult health care providers. The report outlines a decision-making algorithm for managing the transition process. Got Transition, the national center for HCT, developed a structured approach called the Six Core Elements of HCT to help to implement the recommendations of the AAP, AAFP, ACP clinical reports. The Six Core Elements of HCT have been shown to improve HCT process in primary, subspecialty, and Medicaid managed care, as well as children’s hospital clinical settings. Providers can decide which of the core elements to be implemented based on the needs of their patients’ and the practices’ available resources. Implementing a systematic method of
transition for patients with rheumatologic disease is essential in achieving a successful transfer of care; however, the process must be individualized to each patient. Children with rheumatic disease have a wide range of transition needs based on multiple factors such as disease activity, readiness, cognitive and executive functioning proficiency, home support structures, and socioeconomic factors. For instance, a child with mild disease, minimal to no disease morbidity, stable social environment, and excellent accesses to resources will require much less assistance in the transition process. This contrasts with a child who comes from an unstable socioeconomic background and has severe disease with multiorgan involvement who requires multiple medications and management by several subspecialty providers. In this scenario, the rheumatology practice would employ all of the Six Core Element processes.

The Six Core Elements of HCT address the preparation, transfer, and integration phases of transition and are broken down into a transition policy, tracking and monitoring, readiness assessment, transition planning, transfer of care, and transfer completion. Three different packages with tools for each core element are available; for those caring for youth who will transfer to an adult practice, for those who will move to an adult model of care without changing their provider, and for those who will be accepting young adults into their practice. Table 1 shows the Six Core Elements for youth transitioning to an adult health care provider. Figure 1 demonstrates how the Six Core Elements can be applied within a practice setting over the transition planning and transfer process. In this study, each of the elements are discussed from a rheumatological point of view. To assist with implementation of the Six Core Elements of HCT, the ACR joined the ACP Pediatric to Adult Care Transitions Initiative and formed the ACR Transition Work Group. They developed a subspecialty-specific toolkit72 tailored to pediatric and adult rheumatologists to assist them in transitioning their patients.

### Table 1 Contents of the Six Core Elements for “Youth Transitioning to Adult Care”

<table>
<thead>
<tr>
<th>1. Transition policy</th>
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<tbody>
<tr>
<td>• Develop a transition policy/statement with input from youths and families that describes the practice’s approach to transition, including privacy and consent information</td>
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<tr>
<td>• Educate all staff about the practice’s approach to transition, the policy/statement, the Six Core Elements, and distinct roles of the youth, family, and pediatric and adult health care team in the transition process, taking into account cultural preferences</td>
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<tr>
<td>• Post policy publicly and share/discuss with youths and families, beginning at the age of 12–14 years, and regularly review as part of ongoing care</td>
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<tr>
<td>2. Transition tracking and monitoring</td>
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<tr>
<td>• Establish criteria and process for identifying transitioning youths and enter their data into a registry</td>
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<tr>
<td>• Use individual flow sheet or registry to track youth’s transition progress with the Six Core Elements</td>
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<tr>
<td>• Incorporate Six Core Elements into clinical care process, using electronic medical record if possible</td>
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<tr>
<td>3. Transition readiness</td>
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<tr>
<td>• Conduct regular transition readiness assessments, beginning at the age of 14 years, to identify and discuss with youth and parent/caregiver their needs and goals in self-care and how to use the adult health care system</td>
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<tr>
<td>• Jointly develop goals and prioritized actions with youth and parent/caregiver and document regularly in a plan of care</td>
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<tr>
<td>• Develop and regularly update the plan of care, including readiness assessment findings, goals and prioritized actions, medical summary and emergency care plan, and, if needed, a condition fact sheet and legal documents</td>
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<tr>
<td>• Prepare youth and parent/caregiver for adult approach to care at the age of 18 years, including legal changes in decision making and privacy and consent, self-advocacy, and access to information</td>
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<tr>
<td>• Determine need for decision-making supports for youth with intellectual challenges and make referrals to legal resources</td>
<td></td>
</tr>
<tr>
<td>• Plan with youth and parent/caregiver for optimal timing of transfer. If both primary and subspecialty care are involved, discuss optimal timing for each</td>
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<tr>
<td>• Obtain consent from youth/guardian for the release of medical information</td>
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<tr>
<td>• Assist youth in identifying an adult provider and communicate with selected provider about pending transfer of care</td>
<td></td>
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<tr>
<td>• Provide referrals to insurance resources, self-care management information, and culturally appropriate community supports</td>
<td></td>
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<tr>
<td>5. Transfer of care</td>
<td></td>
</tr>
<tr>
<td>• Confirm date of first adult provider appointment</td>
<td></td>
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<tr>
<td>• Transfer young adult when his/her condition is stable</td>
<td></td>
</tr>
<tr>
<td>• Complete transfer package, including final transition readiness assessment, plan of care with transition goals and pending actions, medical summary and emergency care plan, and, if needed, legal documents, condition fact sheet, and additional provider records</td>
<td></td>
</tr>
<tr>
<td>• Prepare letter with transfer package, send to adult practice, and confirm adult practice’s receipt of transfer package</td>
<td></td>
</tr>
<tr>
<td>• Confirm with adult provider the pediatric provider’s responsibility for care until young adult is seen in adult setting</td>
<td></td>
</tr>
<tr>
<td>• Contact young adult and parent/caregiver 3–6 months after last pediatric visit to confirm transfer of responsibilities to adult practice and elicit feedback on experience with transition process</td>
<td></td>
</tr>
<tr>
<td>• Communicate with adult practice confirming completion of transfer and offer consultation assistance, as needed</td>
<td></td>
</tr>
<tr>
<td>• Build ongoing and collaborative partnerships with adult practices</td>
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</tbody>
</table>

**Note:** Data from [www.gottransition.org](http://www.gottransition.org).
Policy
The ACR Transition Work Group recommends that each pediatric rheumatology practice develops a transition policy to share with patients and their families, which can be customized by the provider for their practice. As part of the adult rheumatologist toolkit, the ACR endorses using a patient welcome letter as part of their transition policy. Having a transition policy allows both the youth and the family to understand the transition process in the practice (when it starts, when they are expected to find an adult provider, what support the practice will offer during the process) and know what is expected of them. It can be challenging for providers developing the policy to agree on the practices’ HCT process, decide when and how to share the policy with their patients, and how to ensure all providers follow it. Y/YA and their families state that knowing the process and their roles in it makes transition easier and less stressful for all involved.

Tracking and monitoring
The second element of the Six Core Elements of HCT is establishing a mechanism to track the progress of each individual patient as they advance through the transition process. It is recommended to use an individual flowchart to track each patient’s progress and to use the electronic medical record (EMR) to monitor the transition process of a larger population. For example, a HCT registry was developed during a pilot study by Got Transition and a Washington DC-based managed care plan, Health Services for Children with Special Needs (HSCSN) to assess the effectiveness of customizing and incorporating recommended transition services into a Medicaid managed care plan. The tracking process for individual patients was implemented by using a HSCSN nurse care manager and an AmeriCorps volunteer to reach out to participants, highlighting the importance of a multidisciplinary team to support the transition process. The importance of a team-based approach in tracking patient
progress was also emphasized in feedback interviews for a quality improvement (QI) model to apply systematic clinical transition practices in primary care clinics.22 The QI HCT model effectively implemented the Six Core Elements of HCT and developed written registries to track transitioning youth at all sites. A QI initiative at Texas Children’s Hospital used an EMR-based transition planning tool for youth with SHCN and found that nurses and case managers were the top users,25 underscoring the importance of a team-based approach. Additional efforts to integrate HCT into EMRs to improve patient tracking are needed.

Readiness assessment
A fundamental element of the HCT process is preparing Y/YA to manage their own health, as reported in a recent review.26 This involves the Y/YA, the family, and the provider as equally engaged collaborators. By using a transition readiness assessment, providers can evaluate patients’ self-management abilities. Skills in navigating the health care system and taking responsibility for managing their own health are particularly important for those individuals with chronic illnesses. There have been several validated methods for evaluating transition readiness in patients with various chronic conditions. As part of developing more reliable transition processes, the AAP, AAFP, and ACP guidelines suggest that providers should regularly assess transition readiness by using an objective measure. The tools should be used to learn which skills the youth need to improve their self-management and, for caregivers, to learn which skills they need to help their youth attain. To date, none of the readiness assessment tools have been shown to predict a successful transition.

Several transition readiness assessments were reviewed by completing a literature search under PubMed with combinations of the following terms: transition readiness, assessment, questionnaire, chronic illness, condition, pediatric, and adolescent. Of those reviewed, the readiness assessments discussed below and outlined in Table 2 were most frequently cited for use in children with chronic illness. There are also several non-rheumatology disease-specific readiness assessments that are not discussed here, but reviewed elsewhere.27,28 In addition, we included one rheumatology-specific readiness assessment in our discussion.

1. Got Transition readiness assessment is included in the pediatric rheumatology ACR transition toolkit. The Got Transition readiness assessment scale involves rating transition importance and youth confidence as well as questions regarding patient knowledge of their health and their ability to use and navigate the health care system. Assessing confidence is an important way to evaluate Y/YA engagement in their health, which has been shown to improve health outcomes and is a key issue in this age group.29 The Got Transition readiness assessment was used in several HCT QI projects with successful implementation at all sites.18,22–24 It is available in two forms, one for youth and one for parents, is customizable, available in Spanish, and does not require a provider for administration.1

2. The Transition Readiness Assessment Questionnaire (TRAQ)30 is also included in the ACR toolkit. The TRAQ is a 29-item patient-reported assessment that focuses on two domains: self-management and self-advocacy. It is one of the few available validated, patient-reported, and disease-non-specific transition readiness tools and does not require a provider to administer it. In a study of patients with pediatric-onset chronic health conditions, TRAQ scores were shown to be positively associated with age and typically higher in females.31 Another study reinforces the existing data that young adolescents are often not prepared for transition.15 There have been theorized inadequacies of the TRAQ assessment, including the notion that the TRAQ has a “ceiling effect” in which its utility decreases in individuals with higher scores.32 In addition, it may not be an adequate measure of readiness in younger adolescents.33

3. TRxANSITION Index (formerly TRxANSITION Scale) is a readiness assessment that, unlike the other tools, requires a staff member to administer. The assessment is performed via semistructured interviews, and responses are verified by referring to the patients’ medical records. It comprises 32 items across 10 domains, including Type of chronic health condition, Rx/medications, Adherence, Nutrition, Self-management, Issues of reproduction, Trade/school, Insurance, Ongoing support, and New health providers.34 The STARx (Self-Management and Transition to Adulthood with Rx = Treatment) is a self-reported questionnaire developed to gather information on self-management and HCT skills in youth with chronic health conditions and is complementary to the TRxANSITION Index. It has 18 questions that cover three domains and has been internally validated.35,36
<table>
<thead>
<tr>
<th>Assessment</th>
<th>Validated?</th>
<th>How many questions?</th>
<th>Who administers?</th>
<th>Content</th>
<th>Customizable?</th>
</tr>
</thead>
<tbody>
<tr>
<td>Got Transition: Transitional Readiness Assessment</td>
<td>No</td>
<td>25 questions</td>
<td>Self-administered; parent and patient forms</td>
<td>Patient knowledge of health ability to navigate system</td>
<td>Yes</td>
</tr>
<tr>
<td>Transition Readiness Assessment Questionnaire (TRAQ)</td>
<td>Yes</td>
<td>29 questions</td>
<td>Patient self-administers</td>
<td>Self-management skills self-advocacy</td>
<td>No</td>
</tr>
<tr>
<td>TRxANSITION</td>
<td>Yes</td>
<td>32 questions; semistructured interview format</td>
<td>Administered by clinic staff</td>
<td>Type of chronic health condition Rx/medications Adherence Nutrition Self-management skills Issues of reproduction Trade/school Insurance Ongoing support New health care providers</td>
<td>No</td>
</tr>
<tr>
<td>Self-Management and Transition to Adulthood with Rx = Treatment (STARx)</td>
<td>Yes</td>
<td>18 questions</td>
<td>Patient self-administers</td>
<td>Medication management Provider communication Engagement during appointments Disease knowledge Adult health responsibilities Resource utilization</td>
<td>No</td>
</tr>
<tr>
<td>ON Taking Responsibility for Adolescent/Adult Care (ON TRAC)</td>
<td>Yes</td>
<td>25 questions</td>
<td>Patient self-administers</td>
<td>Self-advocacy and self-esteem Self-management health care behaviors Sexual health Social supports Vocational planning</td>
<td>No</td>
</tr>
<tr>
<td>TRANSITION-Q</td>
<td>Yes</td>
<td>14 questions</td>
<td>Patient self-administers</td>
<td>Self-management skills in health and health care</td>
<td>No</td>
</tr>
<tr>
<td>California Healthy and Ready to Work (CA HRTW)</td>
<td>Yes</td>
<td>90 patient questions 72 parent questions</td>
<td>Self-administered; parent and patient forms</td>
<td>Disease knowledge/management Preventative health behaviors Emergency measures Environmental accommodations Monitoring of health condition Management of health condition Communication Community resources Responsible sexual activity Health records Health insurance Accommodations Developmental needs Transportation</td>
<td>No</td>
</tr>
<tr>
<td>Rheumatology-specific transition readiness skills (RACER)</td>
<td>No</td>
<td>32 questions</td>
<td>Patient self-administers</td>
<td>Planning for adult life Self-advocacy Health service navigation Health condition management</td>
<td>No</td>
</tr>
</tbody>
</table>

questionnaire for assessing transition readiness. Its objective was to provide a transition planning program that was based on self-management. It is divided developmentally into three age-specific transition stages that address six domains of transition: 1) self-advocacy and self-esteem, 2) independent/self-management health care behaviors, 3) sexual health, 4) social supports, 5) educational, vocational, and financial planning, and 6) health and lifestyle. The TRANSITION-Q is a self-administered readiness scale that is used to measure health care management skills. It has been designed using Rasch measurement theory analysis to apply a modern psychometric approach.
of assessing transition readiness, does not focus on other self-management skills such as education or vocation, and has been used in children with different chronic health conditions.

6. The California Healthy and Ready to Work (CA HRTW) Transition Assessment Tool measures the overall transition readiness reported by both patient and parent and is based on 14 domains including patient knowledge of condition, disease management, ability to navigate the health care system, communication, community resources, sexual health, transportation, and developmental needs. Guidelines using HRTW for teaching youth with SHCN health and self-care skills were based on the developmental frameworks of Piaget and Inhelder and Erikson and self-care models of Orem. It has established content validity; however, a pilot study assessing the CA HRTW Transition Assessment Tool found it had low internal consistency for many domains. This may have been attributed to limited range of responses and low samples size. It has also been used in Y/YA with chronic rheumatic disease with data suggesting that older patients were more able to navigate the health care system but did not demonstrate an increased proficiency in medication management.

7. The Readiness for Adult care in Rheumatology (RACER) questionnaire is a self-administered rheumatology-specific assessment. It was developed to measure transition readiness, assess gaps in youths’ knowledge and preparedness, recognize areas in need of additional support, and evaluate efficacy of transition interventions. After consensus agreement, four domains were incorporated into the RACER questionnaire, including 1) planning for adult life, 2) self-advocacy, 3) health service navigation, and 4) health condition management.

Self-management is cited as a central skill needed for transition readiness. Parental support for the development of their youth’s self-management skills and their ability to make independent health care decisions are outlined in a recent literature review by Burke et al. The possession of self-management skills is often dictated by executive functioning proficiency, which continues to develop through the mid-20s. To assess transition readiness, the Kennedy Krieger Independence Scales-Spina Bifida Version was developed as a caregiver-reported measure of self-management skills and executive functioning in youth with spina bifida. Preliminary data support its use as a reliable and valid assessment of self-care management skills. This holds potential value in identifying individual deficiencies in self-management and executive functioning in order to personalize transition planning and readiness assessments.

Resilience, defined as the ability to respond positively to adversity, has been correlated with transition readiness in other chronic conditions. Using the Resilience Scale (RS-14), a validated measure in adolescents and adults, pediatric SLE patients enrolled in a cross-sectional study completed questionnaires to assess whether resilience was associated with transition readiness via TRAQ assessment. Results indicated that TRAQ scores were significantly correlated with resilience and age, but not associated with disease duration, fatigue, anxiety, depressive symptoms, disease damage, or disease activity. Thus, correcting for age and other confounding factors, resilience scores seem to be correlated with transition readiness assessment scores, and it may be important to include resilience building in transition programs.

Regardless of the tool used to assess transition readiness, the AAP, ACP, and AAFP guidelines recommend providers to assess Y/YA and family members independently to determine the level of the Y/YA’s independence.

Transition planning

Transition planning is a collaborative and fluid process which is driven by the Y/YA’s medical problems and individual goals. Got Transition guidelines include examples of a HCT plan of care and suggest to regularly update the plan of care using the readiness assessment findings and to incorporate the goals and prioritized actions of the Y/YA and their families. HCT planning also includes developing a portable medical summary and an emergency care plan that can be sent to the adult providers in advance of the Y/YA’s transfer.

Timing has been shown to be a key component of transition planning. The 2011 AAP, AAFP, and ACP clinical reports recommend that initiating the first step in the transition planning process should occur at the age of 12 years and that transfer to an adult provider should occur between the ages of 18 and 21 years. Further emphasizing the importance of timing, the AAP, ACP, and AAFP clinical recommendations include an algorithm with guidelines for implementing specific stages of transition planning based on the most appropriate age. Within these recommendations, the optimal time of transfer should be individualized to the needs of each Y/YA and all primary and subspecialty care providers should be involved.

Transition planning also includes preparing youth and the parent/caregiver for legal changes in decision making, if needed, and should include obtaining necessary
legal documents as the youth approaches the legal age of adulthood. For some youths and families, this may require referring them to legal resources about supported decision making, and for all young adults, it may require obtaining their consent to involve parents/caregivers in their care when they are legally an adult.71

Finally, transition planning involves inquiring about youth’s preferences for an adult provider and assisting them in the process of finding one. A current and vetted list of adult primary and specialty care providers interested in caring for young adults should be shared with youths and families. Patients should be given information regarding insurance resources, self-care management, and culturally appropriate community supports. The key to successful transition planning is regularly updating the youth’s plan of care. Transition planning can be as simple as the next steps with pertinent goals documented in a clinic note, or more involved with detailed plans of care for Y/YA with complex medical problems.

Transfer/communication

The AAP, ACP, and AAFP clinical reports emphasize the importance of communication between pediatric and adult providers for the exchange of medical and special non-medical information.28 Sharing of a medical summary along with a transfer letter with the adult provider is a core component of the ACR toolkit for pediatric providers. The transfer letter provides a brief introduction to the patient and includes personal details that will help to foster the patient–adult provider relationship. Medical summary templates for patients with JIA and SLE are available in the ACR toolkit.

There is an increasing evidence supporting the value of checklists for successful transfer of care. Checklists have long been utilized in other areas of medicine, such as surgery, with significant improvements in patient outcomes as evidenced by reductions in surgical complications and deaths.50 In adults transitioning from hospital to home, interventions that include a structured transition program with a guide or checklist have been associated with decreased rehospitalizations and medical costs.51 There has been a growing recognition that they have been associated with decreased rehospitalizations and medical costs.51 There has been a growing recognition

This Delphi study defined what themes, specific to youth with chronic rheumatologic disease, should be addressed during the process of transition.52

Transition completion/feedback

Obtaining Y/YA feedback is an important step in the transition process; an example of a transition feedback survey is included in the Six Core Elements packages. In a pilot study implementing the Six Core Elements of HCT, transition feedback survey results of 10 pilot group members revealed inadequacies in the discussions of privacy and consent as well as in provider help in gaining self-care skills.18 In addition, the creation and distribution of a medical summary were reported in only half of the respondents. Another feedback survey was used in youth with JIA is the Mind the Gap scale that assessed satisfaction with transition to an adult provider in 10 major UK rheumatology centers.53 There are several other feedback surveys that can be used in Y/YA with many different chronic conditions to assess the preparation for transfer and transfer experience. The ADolescent Assessment of Preparation for Transition is both a reliable and a valid survey for adolescents with chronic health conditions between the ages of 16 and 17 years. It comprises 26 items including self-management, medication management, and planning for transfer of care.54 The On Your Own Feet Transfer Experience Scale (OYOF-TES) consists of 18 items rated on 5-point Likert scales. Research using the OYOF-TES showed good reliability and internal consistency and underscored the importance of collaboration between pediatric and adult providers.55 The Patient Activation Measure (PAM) for Adolescents, initially tested in a large sample of adolescents with various chronic healthy conditions, measures patient engagement of adolescents with chronic conditions among a broad age range. It has good test–retest reliability and internal consistency.56 Continued analysis of patient satisfaction of their transition process and outcome will provide valuable feedback on what elements should be included in transitional care programs.

Role of the adult provider in transition

HCT does not end with transfer of the patients. Smooth integration of patients into adult practices can be one of the more difficult aspects of HCT. Usually, more than one adult appointment is required to establish and nurture the new patient–provider relationship. During the transition process, providing support for and emphasizing the role of the adult provider have been shown to be key indicators for successful transition.57 By using Delphi analysis, Suris and Akre found that 50% of the indicators for successful transition
concentrated on engagement between the patient and the adult provider, further emphasizing the importance of the role of adult providers. To assist with adult rheumatologists’ participation in the transition process, the ACR toolkit provides handouts specific to the adult providers and offers examples of a welcome letter.

**Transition models**

The Six Core Elements of HCT is a process that can be incorporated into any model of care and is not disease-specific. There have been several proposed models of transition care within the field of rheumatology; however, to date, no model has been shown to result in measurable improved outcomes. In the UK, McDonagh et al proposed a transition program based on needs assessments of focus groups, a national survey of health professionals, Delphi analysis, and retrospective case audits. In the analysis of the impact of the program on adolescents with JIA, significant improvements were observed in health-related quality of life and vocational preparation. However, it is unknown if these observations could be attributed to participation in the transitional care program or reflected improvement in disease activity or patient maturation. In addition, it is unknown whether these improvements persist into adulthood or if they are associated with positive long-term outcomes. A transition model for young adults with rheumatic disease, described by Tucker and Cabral, proposed a shared clinic between pediatric and adult rheumatologists, with assistance of a multidisciplinary team. This clinic encouraged patient independence while discouraging parental involvement. Another rheumatology-specific transition model that uses a multidisciplinary team was described by Rettig and Athreya, which uses a nurse coordinator. The program comprises pre-transitional assessments and interventions that address education, vocation, and sexuality.

To facilitate the existing models of transition of care for young people with juvenile onset of rheumatic and musculoskeletal disease, a task force of the Pediatric Rheumatology European Society (PRES)/European League Against Rheumatism (EULAR) developed the first international set of recommendations and standards. It includes 12 consensus recommendations of “essential/minimal” and “ideal/optional” components of transitional care established via international Delphi analysis and systemic literature review. The recommendations are synchronous with the 2016 National Institute for Health and Care Excellence (NICE) guidance on Transition in the UK. The NICE guidelines offer recommendations that are characterized by youth involvement and partnership in the transition process and emphasize developmentally appropriate transition support as well as a strength-based practice that focuses on a young person’s individual skills and experience to achieve the intended outcomes.

**Evidence supporting implementation**

There is a growing evidence that a structured transition program improves outcomes, and there is evidence supporting the feasibility of implementing structured transition programs. Recently, the University of Rochester Medical Center integrated the Six Core Elements into their clinical practice and saw the percentage of patients tracked through transfer of care more than double compared to baseline. Pediatric providers successfully implemented the Six Core Elements of HCT by using transition readiness assessments and improved communication with adult providers through medical summaries and closed-loop exchanges. There has also been a recent evidence in workshops to increase transition skills in adult rheumatology providers. Combining objective standardized clinical examination sessions with skill-based transition workshops has shown to increase adult rheumatology fellows’ confidence in transition skills and ability to employ best transition practices. Providing curriculum-based training to all rheumatologists-in-training is a system-based approach that may improve provider knowledge and confidence around transferring patients from pediatric to adult care.

**Measuring process and outcomes**

Many of the readiness assessments and transfer recommendations have not yet been validated against measurable outcomes of successful transition. Efforts to accomplish outcome-related research have failed to fully address the transition needs of adolescents. The general lack of specific outcome measures has been cited as problematic, as reported in a recent review of transition of care programs performed by the EULAR/PRES working group. There have been several proposed quantifiable outcomes to measure successful transition including uninterrupted access to health care, improved follow-up rates of patients transferred, and less “no show” visits. Huang et al studied the capability of a generic technology platform to improve disease management tasks and health-related self-efficacy in a transition cohort of patients with type 1 diabetes, inflammatory bowel disease, and cystic fibrosis and found improvement in disease management as a short-term outcome.
studies of transition outcomes in childhood onset of SLE, Son et al showed that lower educational level (less than a high school degree) was associated with increased missed appointments in the posttransition period, indicating that at-risk populations may benefit from additional guidance during the transition process. Recently, a UK study investigated the clinical impact of the DON’T RETARD project (Devices for Optimization of Transfer and Transition of Adolescents with Rheumatic Disorders). The DON’T RETARD transition program included a transition coordinator and provided information about JIA and medication management as well as how to manage fatigue, friendships, and schoolwork. Its effect on primary clinical outcomes including physical, psychosocial, and rheumatic-specific health status and secondary outcomes of improved quality of life were assessed in youth with JIA. Results indicated that implementation of a transition program as a brief intervention can improve the perceived health and quality of life of adolescents with JIA during the transition process. Furthermore, the study showed that a randomized controlled trial can be designed to evaluate the effectiveness of a transition program. Unfortunately, the study only assessed clinical differences rather than statistical significances, was performed at a single center, and did not assess long-term outcomes. Another UK study assessing transition outcomes after implementation of an evidence-based transitional program reported improvements in health-related quality of life, knowledge, satisfaction, and vocational readiness markers.

Investigators have begun assessing the facilitation into employment of Y/YA with childhood onset of rheumatic and musculoskeletal diseases as a long-term outcome measure. There are several avenues of provider-directed guidance that patients have identified as helpful, such as access to flexible and convenient care, information and policy sign-posting, social and emotional support, skill training, and advocacy. Studies indicate that emphasis should be placed on provider proficiency in informational assistance and advocacy as there is often a lack of knowledge regarding antidiscrimination legislation. In addition, to improve employment outcomes, patients need support in disclosing their disease and its physical impact on their employment, which underscores the necessity of building their social skills and offering emotional support in a HCT program through a multidisciplinary team. Anxiety regarding disease disclosure is not disease-specific and interventions such as role play and vocational readiness counseling programs have been positively evaluated in other specialties. These avenues of provider-directed guidance should be supported by a strong foundation of optimal disease management. The attainment of successful employment following these interventions has not been adequately described.

An approach to assess and categorize transition measures can be done using the “Triple Aim” conceptual model. The Triple Aim was developed by the Institute for Health Care Improvement and consists of three interdependent goals: 1) to improve the individual experience of health care, 2) to improve the health of populations, and 3) to reduce the per capita costs of care. A systematic review used the Triple Aim model to categorize measures evaluating transition interventions in studies of transition of adolescents with and without chronic conditions. In the review, patient-reported health outcomes such as social and physical functioning, pain, mental health, vitality, emotional and physical roles, general perceived health, and quality of life were measured by using validated surveys. Other outcomes measured include adherence to care, follow through of medication and screening recommendations, fluctuations in drug levels, and patient mortality. By using the Triple Aim model to assess transition of care studies, the authors concluded that transition intervention studies measure the impact on population health, patient experience, and cost inconsistently. Thus, more consistent measures need to be implemented to better assess patient experience and outcomes. More recently, provider experience has been added, and it is now called the Quadruple Aim to measure the outcomes.

Conclusion and recommendations

to adopt a consensus on outcome measures and a better understanding of who is at risk for poor outcomes. All youths require transition support; however, identifying the factors that put particular youth more at risk for a poor transition outcome would allow the correct amount of resources to be targeted for the appropriate population. Offering appropriate individualized patient support will likely improve some of the outcomes of Quadruple Aim. There is currently a lack of rheumatology-specific data that focuses on measuring the outcomes of Quadruple Aim. There is currently a lack of rheumatology-specific data that focuses on measuring the outcomes of Quadruple Aim, and further studies are needed regarding the improvement of patient and provider experiences and reduction of health care costs. The next steps of research in transition of care should include assessing different transition care models, using a common process of preparation, transfer, and integration, and incorporating the youth’s level of needed HCT supports. In addition, further studies to assess how aspects of each model affect
the outcomes are needed. Transition model components such as care coordinators, team models, combined clinics, and young adult clinics should be more closely evaluated. In addition, disease-specific models have been the focus of much research, the necessity of which is unknown. Finally, improving payment models for transition of care in countries that do not have publicly funded health care would assist in the implementation of transition processes. Current fee-for-service payment strategies with vignettes are outlined in a Got Transition Payment and Reimbursement Tip Sheet, but a more comprehensive approach beyond fee-for-service needs to be studied and best practices implemented. Although much progress has been made in understanding the barriers to transition, valid readiness assessments and the feasibility of implementing different models of transition programs, there are many unanswered questions. Future directions should focus on how to pay for transition of care in countries where important, which outcomes should be measured, how best to measure them, who is at risk for poor outcomes, and what transition model components are most significant to improve outcomes.

Disclosure
The authors report no conflicts of interest in this work.

References


