Primary care interventions to improve transition of youth with chronic health conditions from paediatric to adult healthcare: a systematic review

Jasmin Bhawra,1 Alene Toulany,2,3,4 Eyal Cohen,1,4,5 Charlotte Moore Hepburn,1,4 Astrid Guttmann1,3,4,5

ABSTRACT

Objective: To determine effective interventions to improve primary care provider involvement in transitioning youth with chronic conditions from paediatric to adult care.

Design: Systematic review. Multiple electronic databases were searched including Ovid MEDLINE, EMBASE and Web of Science (from 1 January 1947 to 5 August 2015). Evidence quality was assessed using a 36-point scoring system for disparate study designs.

Setting: Studies with paediatric-to-adult transition programmes and interventions involving primary care providers or in primary care settings.

Participants: Youth aged 16 years and over.

Outcomes: Relevant outcomes were grouped into 3 main domains based on the Triple Aim Framework: experience of care, population health, cost.

Results: A total of 1888 unique citations were identified, yielding 3 studies for inclusion. Overall, primary care provider roles were not well defined. 2 studies used case managers to facilitate referrals to primary care, and the remaining study was the only 1 situated in a primary care setting. None of the studies examined transition in all 3 Triple Aim Framework domains. The most commonly reported outcomes were in the cost domain.

Conclusions: There is limited empiric evidence to guide primary care interventions to improve transition outcomes for youth with chronic conditions. Future research and policy should focus on developing and evaluating coordinated transition interventions to better integrate primary care for high need populations.

INTRODUCTION

An estimated 15–18% of youth in North America have a chronic health condition and over 98% of these youth are expected to reach age 20, thereby requiring transition from paediatric to adult healthcare.1 2 Although there is no universally accepted age of transfer (one-time event), many jurisdictions use inflexible age cut-offs to delineate service boundaries, creating fragmentation and discontinuity in provision of care across a particularly vulnerable period.3 4 The transition period encompasses multiple steps in a process including thoughtful planning, the actual transfer from paediatric to adult care, and adjustment to the new system afterwards. As such, it can be challenging for patients, families and clinicians alike.5 6 More than half of the youth with chronic health conditions report inadequately support and services during their transition to adult healthcare.7–9 It is well recognised that the process is often flawed due to inadequate planning, poor service coordination, lack of resources and gaps in

Strengths and limitations of this study

▪ This systematic review used a rigorous methodology and was an exhaustive search of all studies of transition interventions involving primary care with no restrictions placed on language, jurisdiction, or disease type.

▪ The main limitation of this systematic review is the small number of studies (three) that have assessed primary care-based transition interventions even with inclusion criteria that did not mandate randomised trial design. In addition, differences in primary care provision for children across various healthcare jurisdiction makes generalisability of study findings challenging. The only study that was based in a primary care setting was in paediatrician-led US practices which may not be relevant to systems where general practitioners provide ‘cradle to grave’ primary care.

▪ The review highlights the need for more rigorous intervention studies to guide improved integration of primary and specialised care during transition and better define the role of primary care providers in facilitating an effective transition for youth with chronic conditions.
education and training. Transition for youth with chronic health conditions is also complicated by the intricate physical and psychosocial developmental changes that accompany adolescence. Together, these challenges contribute to potentially serious health-related consequences including patient disengagement, poor treatment adherence, increased hospitalisation rates and overall detrimental health outcomes.

The goal of healthcare transition for youth with chronic health conditions should be to maximise long-term functioning through the delivery of high-quality, developmentally appropriate healthcare services that continue uninterrupted during the passage from adolescence to adulthood. To date, the majority of programmes designed to enhance transition for youth with chronic illness have been condition-specific or subspecialty-specific and hospital-based with little focus on primary care. Much of the currently available literature is primarily descriptive, focuses on the transfer component (one-time event) of transition (purposeful, planned process) and is limited to only a few studies measuring the effectiveness of any particular transition intervention.

Although national bodies have recommended more effective primary care provider involvement during transition and integrated primary and specialist care, the role of primary care in facilitating transition and potentially mitigating adverse events associated with changes in care for youth with chronic conditions remains unclear. In addition to specialty care, paediatric to adult primary care transition may also be needed for youth in health systems that rely heavily on paediatricians for primary care. Few transition programmes base their services in primary care settings and utilise the primary care provider to assess and coordinate the health needs of the adolescent and family. The aim of this systematic review is to determine effective interventions to improve the role of primary care providers in the transition of youth with chronic health conditions from paediatric to adult healthcare.

METHODS

Search strategy

A systematic review of all literature and conference proceedings published in the past 65 years (1 January 1947 to 5 August 2015) was conducted using EMBASE, Web of Science, Ovid ‘MEDLINE’ and Ovid ‘In Process & Other Non-indexed Citations’. Authors of all relevant conference abstracts were contacted to enquire about publications, as only published articles are included in the final results.

The literature search focused on interventions and evaluations targeting a paediatric population (using terms such as ‘youth’, ‘adolescent’ and ‘paediatric’), transition (ie, ‘continuity of care’ and ‘transfer’) and primary care (ie, ‘medical home’, ‘family practice’, ‘family physician’ and ‘paediatrician’). Articles published in all languages and all study designs were included in the search. The search was not limited to specific diseases in order to be inclusive of all chronic medical and mental health conditions (see online supplementary file for full search strategy). Articles discussing opinion-based best practice, those defining family/caregivers as primary care providers and transfers between hospital services or departments were excluded.

Study selection

Two reviewers (JB and AT) screened all article titles and abstracts. The articles were first screened independently and then examined in full by both reviewers for the inclusion criteria. All identified non-English papers were in French and were assessed by a bilingual member of the research team (AG). The reviewers met to discuss any disagreements which were resolved by consensus (JB, AT, EC and AG).

Data synthesis and quality analysis

Data from the final included studies were evaluated using a data abstraction table summarising study characteristics, level of primary care engagement and outcomes as per the Triple Aim Framework domains (experience of care, population health, cost). The reviewers identified which Triple Aim domains were assessed by each study outcome (table 1). Quality scores were also calculated for each of the articles using Hawker et al’s scoring system for disparate study designs. A 36-point scale based on nine domains (abstract and title, introduction and aims, methods and data, sampling, data analysis, ethics and bias, results, transferability or generalisability, and implications and usefulness) was used to grade each study. Two authors (JB and AT) scored the studies and resolved disputes by consensus.

RESULTS

Literature overview

Figure 1 illustrates the results of the search and screening strategies. A total of 1888 unique citations were

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Triple Aim transition intervention measures</th>
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<tbody>
<tr>
<td><strong>Experience of care</strong></td>
<td><strong>Population health</strong></td>
</tr>
<tr>
<td>◀ Satisfaction</td>
<td>▶ Adherence to care/guidelines</td>
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<tr>
<td>◀ Barriers to care</td>
<td>▶ Disease-specific measures</td>
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<td>▶ Mortality</td>
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<td>▶ Patient-reported outcome measures</td>
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<td>▶ Process of care</td>
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<td>▶ Self-care skills</td>
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Adapted from Prior et al30 and Stiefel and Nolan.31
identified and screened. In total, 1061 articles were excluded at the abstract screening stage and 45 citations (including 27 journal articles and 16 conference abstracts) were assessed for eligibility. Twenty-seven citations were excluded after full-text review because they did not discuss transition in a primary care setting. Ten of these citations were relevant commentaries on the role of primary care in transition, but were not interventions or evaluations. In total, seven citations discussed transition interventions or evaluations and primary care, including three journal articles and four conference proceedings, of which only one was eligible. The authors of the abstracts were contacted; however, there were no associated article publications of the preliminary results.

Study characteristics
The three included studies selected were conducted in the USA (n=2) and Canada (n=1). The studies sampled youth with chronic health conditions, sickle cell disease (SCD), and type 1 diabetes. The programme/intervention target groups were most often patients and their families; however, feedback from healthcare providers was an outcome in one of the three studies. There were no randomised controlled trials in primary care settings or with primary care providers. The three articles included one retrospective cohort, one uncontrolled cohort and a one-time series comparative study. Table 2 describes each study by intervention/programme, patient condition, location, primary care engagement strategy, study design, outcomes/results and Triple Aim Framework domain addressed. Calculated quality scores for all studies yielded a mean of 26.67 (SD 3.33) and median of 25 (range 24.5–30.5). The assessment of risk of bias in individual studies was assessed systematically at the study and outcome levels using the quality score framework described by Hawker et al for disparate study designs.

Studies
Hankins et al conducted a retrospective cohort study of a pilot programme aimed at helping adolescents aged 17–19 years with SCD to find an adult medical home. The study assessed 83 adolescents’ participation, patient/family/provider acceptance of the programme and number of appointments with adult haematologists fulfilled as an indication of programme efficacy. The programme entailed a tour of adult SCD programmes for patients and families and a lunch with programme staff to discuss transition issues such as expectations, barriers and insurance. A paediatric haematology nurse case manager also scheduled patients’ first visit with an adult SCD programme or healthcare provider. This study measured participants’ versus non-participants’ attendance, completion of a first appointment with an adult healthcare provider, as well as patient/parent/provider satisfaction with the programme using surveys. Primary care was involved to the extent that a case manager helped establish patients’ relationship with an adult medical home or healthcare provider.

McManus et al’s time series comparative study involved five health centres which adopted the American Academy of Paediatrics’ (AAP) Six Core Elements of Health Care Transition to assess
### Table 2  Summary of findings

<table>
<thead>
<tr>
<th>Transition intervention/programme</th>
<th>Patient condition</th>
<th>Location</th>
<th>Primary care engagement</th>
<th>Study design</th>
<th>Outcomes/results</th>
<th>Quality score</th>
<th>Experience of care</th>
<th>Population health</th>
<th>Cost</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pilot programme aimed at improving continuity of care by helping adolescents identify an adult medical home. Consists of 3 components including a tour of SCD programmes, lunch discussion with staff, and team scheduling patient’s 1st visit to the adult programme. Attendance records consulted and patient/parent and healthcare provider feedback obtained from surveys.</td>
<td>SCD</td>
<td>Tennessee, USA</td>
<td>Case manager facilitated 1st appointment to help establish relationship with adult medical home/provider of patients’ choice.</td>
<td>Retrospective cohort study</td>
<td>Overall participation ▶ 34 of the 83 agreed to participate (41%) □ ▶ Proportion of participants who fulfilled 1st appointment with an adult haematologist (74%) vs non-participants (33%) in 3 months □</td>
<td>25</td>
<td>X</td>
<td>X</td>
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<td>5 Academic paediatric and adult health centre teams adopted a 2-year learning collaborative to implement ‘Six Core elements of Healthcare Transition’. Teams consisted of a physician and a transition care coordinator and utilised the Health Care Transition Index to assess programme progress in Chronic physical, develop-mental and mental health conditions</td>
<td>Chronically ill</td>
<td>District of Columbia, USA</td>
<td>Study set in 5 large primary care practices. 2 Were in adolescent clinics, 1 in a paediatric clinic, 1 in a family medicine resident clinic, and 1 in an internal medicine clinic.</td>
<td>Time series comparative study(compared results of transition index at 3 points in time over a 22-month period)</td>
<td>Improvements in 6 quality indicators of transition: ▶ Development of an office transition policy (1) ○ ▶ Staff and provider knowledge and skills related to transition (1) ○ ▶ Identification of transitioning youth registry (1) ○</td>
<td>30.5</td>
<td>X</td>
<td>X</td>
<td></td>
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<tr>
<th>Transition intervention/programme</th>
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<th>Cost</th>
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<tr>
<td>implementing the Six Core Elements.</td>
<td>Type 1 diabetes</td>
<td>Manitoba, Canada</td>
<td>A navigator facilitated referrals connecting patients to family physicians and other support when requested.</td>
<td>Uncontrolled cohort study</td>
<td>▶ Transition preparation of youth (1) ◊</td>
<td>24.5</td>
<td>X</td>
<td></td>
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<tr>
<td>A feasibility and acceptability study of the Maestro Project—community-based administrative support and systems navigation service. Patients were contacted biannually to enquire about access to care, services, health status associated with diabetes complications and offered follow-up support.</td>
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SCD, sickle cell disease.
improvement in the development of a transition policy, transitioning youth registry, transition preparedness, transition planning, transfer of care and transition documentation. The study was set in primary care centres, each of which participated in a 2-year learning collaborative and had a team consisting of a lead physician, care coordinator focused on transition (ie, nurse, social worker) and consumer (parent/caregiver or young adult). Teams attended five 1½-day learning and coaching sessions. Sites were provided with financial support to cover staff time and patient participation. Each health centre evaluated their practice on the Core Elements using the Health Care Transition Index, and rated themselves in the six categories using a combination of narrative descriptions and raw scoring. The evaluation was completed three times over the course of the study.

The Maestro Project based in Manitoba, Canada is a navigation service to help young adults aged 18–30 years with type 1 diabetes transfer to adult healthcare services. The Maestro Project maintains a database of 473 young adults living in Manitoba, north-western Ontario, and south-eastern Saskatchewan. The term ‘Maestro’ refers to an administrative project coordinator who provides support (via phone, email, newsletters or drop-in group sessions), identifies barriers to accessing healthcare services and connects young adults with specialty care (ie, general internists, adult endocrinologists), diabetes educators and counsellors, and family physicians. The Maestro also works with community centres involved in diabetes education in order to provide referrals and facilitate follow-up visits. This uncontrolled cohort study assessed the feasibility and acceptability of the Maestro Project by collecting participants’ demographic information, self-reported medical outcomes at baseline and follow-ups, programme participation, project referrals and community connections using surveys administered biannually. Appropriate primary care providers were contacted by the Maestro to support diabetes care for youth transitioning to adult services.

**Triple Aim Framework domains and study outcomes**

**Experience of care**

Experience of care was only assessed by the SCD retrospective cohort study, which used satisfaction surveys administered to patients, parents and healthcare providers. Hankins et al. reported that all participants, parents and healthcare providers found the programme to be either ‘helpful’ or ‘very helpful’. None of the other studies reviewed collected data on patient satisfaction or barriers to care.

**Population health**

Population health measures were used in two of the three studies reviewed. The McManus et al. time series comparative study of five health centres measured population health domain as adherence to care guidelines (using the AAP Clinical Report on Supporting the health care transition from adolescent to adulthood in the medical home) and found improvements in all six transition quality indicators.

**Cost**

Health service utilisation (participation/attendance) was measured in all studies. In the SCD retrospective cohort study, 41% of youth agreed to participate in the pilot programme, and 74% of participants completed their first appointment with an adult haematologist (compared with 33% of non-participants). Similarly, there was an increase in project referrals and community connections among Maestro Project participants. McManus et al. reported improvements in transfer of care scores after implementing the learning collaborative in participating health centres over 15 months. No studies assessed actual costs either as an outcome or an input associated with a transition programme or intervention.

**DISCUSSION**

This systematic review demonstrates that there is little empiric evidence to guide clinical practice or policy around the role of primary care providers in the transition of youth with chronic health conditions. We found only three published studies that evaluated any kind of primary care-specific intervention aimed at improving transition, only one of which was based in a primary care setting. The level of primary care engagement was limited. Two of the interventions included a case manager who facilitated appointments with a primary care provider during transition and therefore did not focus on primary care as the key target of intervention. Overall, none of the studies examined transition measures across all three Triple Aim Framework domains.

In a previous work, we have shown that very few jurisdictions have any health system strategy to address transition. However, in a joint clinical report published by the AAP, American Academy of Family Physicians (AAFP) and American College of Physicians (ACP), primary care providers and medical specialists were encouraged to adopt a transition planning algorithm for all youth within a medical home specific to their clinical setting. However, much of the emphasis in the transition literature (which most likely reflects clinical practice) focuses solely on specialist-to-specialist provider transition. This may reflect in part the de facto interest and leadership of the paediatric specialist community, as well as the generally fragmented nature of the primary to specialty care interface.

There is no gold standard for the organisation of care that will best meet transition needs for all youth with chronic conditions. A range of system-level approaches are most likely needed to improve the process. The level and type of primary care involvement during transition should differ depending on the condition. For example, for some diseases such as uncomplicated...
epilepsy, the primary care provider may be the appropriate centre of medical management across transition, whereas for other more complex conditions such as cystic fibrosis, ongoing management will be centred at a specialised adult centre with primary care involved in a more supportive role. Youth with medical complexity may require a mixed strategy as they often have multiple specialist providers, but often care coordination occurs either in general paediatric clinics in children’s hospitals, or by primary care paediatricians in a medical home.25 28 51 Many paediatric chronic diseases, however, may benefit from a shared-care model or a more staged approach where there is a phased transfer between paediatric subspecialists and primary care providers in a disease-specific way; however, we could not find any evidence to guide these types of approaches.

For any health system strategy around transition, primary care models for children will be an important consideration. In the UK, Australia and most parts of Canada, youth receive primary care from a general or family practitioner and are likely to continue with the same primary care provider as they move into adulthood. Such patients may still benefit from a process to help them improve readiness and increase responsibility for their care as they transition.1 6 Other youth with chronic conditions whose primary care provider is a paediatrician, as opposed to a general practitioner, may experience a ‘double hit’ as they require transition from both primary and secondary or specialist care. We found no data to guide these ‘double hit’ transitions and whether they should occur simultaneously or in a sequential fashion. It is also striking that although the most common chronic conditions of childhood (eg, asthma) are managed almost exclusively by primary care providers, there are no studies evaluating primary care transition models in these populations in systems that rely heavily on paediatrician-led primary care, such as in most states in the USA.

CONCLUSION

This systematic review is limited by the lack of substantial evidence on the role of primary care in enhancing the quality of transition care for youth with chronic conditions. System strategies that account for differences in paediatric primary care providers and the levels of complexity of chronic conditions will be important. Other policy initiatives that seek to improve coordination of care, such as Accountable Care Organizations in the USA, may provide an opportunity to address the role of primary care in transition for youth across transition. It will be essential to establish consistent goals for primary care in transition in order to build an adequate body of literature to affect policy and practice. Future research and policy should focus on developing and evaluating coordinated transition interventions involving primary care to better integrate primary and specialised care.

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