Transition from child to adult health services for young people with cerebral palsy in Ireland: a mixed-methods study protocol


Published in:
BMJ Open

Document Version:
Publisher's PDF, also known as Version of record

Queen's University Belfast - Research Portal:
Link to publication record in Queen's University Belfast Research Portal

Publisher rights
Copyright 2020 the authors.
This is an open access Creative Commons Attribution-NonCommercial License (https://creativecommons.org/licenses/by-nc/4.0/), which permits use, distribution and reproduction for non-commercial purposes, provided the author and source are cited

General rights
Copyright for the publications made accessible via the Queen's University Belfast Research Portal is retained by the author(s) and / or other copyright owners and it is a condition of accessing these publications that users recognise and abide by the legal requirements associated with these rights.

Take down policy
The Research Portal is Queen's institutional repository that provides access to Queen's research output. Every effort has been made to ensure that content in the Research Portal does not infringe any person's rights, or applicable UK laws. If you discover content in the Research Portal that you believe breaches copyright or violates any law, please contact openaccess@qub.ac.uk.
ABSTRACT

Introduction The transition from child to adult health services is a challenging and complex process for young people with cerebral palsy (CP). Poorly managed transition is associated with deterioration in health, increased hospitalisations and reduced quality of life. While international research identifies key practices that can improve the experience and outcomes of transition, there is a paucity of data in the Irish context. This research study aims to gain an insight into the experience of transition for young people with CP in Ireland.

Methods and analysis A convergent parallel mixed-methods design will be used to collect, analyse and interpret quantitative and qualitative data. Participants will be young people aged 16–22 years with CP, their parent(s)/carer(s) and service providers. Quantitative and qualitative data will be collected through questionnaires and interviews, respectively. Quantitative data will be reported using descriptive statistics. Where sufficient data are collected, we will examine associations between the experience of transition practices and sociodemographic and CP-related factors, respectively, using appropriate regression models. Associations between service provider characteristics and provision of key transition practices may also be explored using appropriate regression models. Qualitative data will be analysed using the Framework Method. A coding matrix based on key transitional practices identified from the literature will be used to identify convergence and divergence across study components at the integration stage.

Ethics and dissemination The study has been approved by the RCSI University of Medicine and Health Sciences Research Ethics Committee (REC201911010). Results will be presented to non-academic stakeholders through a variety of knowledge translation activities. Results will be published in open access, peer-reviewed journals and presented at national and international scientific conferences.

INTRODUCTION

Cerebral palsy (CP) is one of the most prevalent disabling conditions among children worldwide. CP is an umbrella term used to describe a group of lifelong conditions that affect movement and posture causing activity limitations, which result from disturbances to the developing fetal or infant brain. Many people with CP also experience associated impairments such as sensory, communication and cognitive impairments. The type and severity of impairment vary considerably among people with CP. The majority of children with CP will survive into adulthood, with many having a life expectancy similar to that of the general population. Continual improvements in longevity have resulted in a transitional care period between child and adult health services for young people with CP. Transition is defined as ‘the purposeful planned process that addresses the medical, psychosocial, educational and vocational needs of adolescents and young adults with chronic medical and physical conditions as they move from child-centred to adult-orientated healthcare systems’. Despite the
importance of transition for supporting the long-term health and well-being of people with chronic conditions, international research indicates that this process is poorly managed for young people with CP. Poor management of transition is associated with deterioration in health, increased hospitalisations and reduced quality of life among young people with long-term conditions. For young people with cerebral palsy, transition from child to adult health services coincides with a decrease in visits to specialist and coordinated services, difficulties accessing clinical care and an increase in unmet health needs. This is concerning given a growing body of evidence shows that adults with CP experience an increasing number of physical and mental health conditions with age, such as cardiovascular disease, arthritis, depression and anxiety. These conditions often co-occur with worsening physical functioning, pain, fatigue and joint deformities due to musculoskeletal progression, or the comorbidities associated with CP. Poor management of transition from child to adult health services may therefore contribute to the development of chronic conditions among adults with CP.

A recent review of the quality of care provided to young people with CP in the UK highlighted that young people have difficulty in navigating adult services, a lack of multidisciplinary teams and leads for adult neurodisability care, and that young people increasingly rely on general practitioners (GPs), after discharge from child services. Further challenges to implementing successful transition for young people in the UK include differences between the funding and organisational structure of child and adult services, lack of service level coordination, lack of involvement of GPs in transition planning and inadequate training of adult health professionals in paediatric conditions. However, a recent programme of research also identifies key practices associated with improved outcomes including a level of parental involvement that is deemed appropriate by both the young person and parent, promotion of health self-efficacy, meeting the adult team before transfer and incorporation of developmentally appropriate healthcare throughout the transition process.

**Transition in an Irish context**

The National Policy and Strategy for the Provision of Neuro-Rehabilitation Services in Ireland and the Integrated Care Programme for Children emphasise the need for appropriate planning and clear protocols to support transition to adult services for young people in Ireland. However, there is no research examining transition for young people with CP in Ireland. Two studies have examined the experience of transition in an Irish context by surveying parents of young people with cystic fibrosis and consultant psychiatrists working in mental health services, respectively. Both studies identified a lack of standardised national practice, lack of written transition plans, lack of interaction between child and adult services and lack of information on self-management and signposting to alternative supports. While this research suggests that transition is poorly managed within the Irish healthcare system, examination of whether these findings are applicable to young people with CP warrants investigation. Moreover, existing evidence regarding transition in the Irish context focuses on the views of parents and service providers. The voice of young people with disabilities or long-term conditions has not been engaged to date.

In order to facilitate implementation of successful transition for young people with CP in Ireland, we need to first assess the gaps in the current management of transition, and second explore the experiences of transition from the multiple perspectives of young people with CP, their families and health professionals on how to implement successful transition in an Irish context.

**Aim**

The aim of this study is to examine how transition care is provided to and experienced by young people with CP in Ireland.

The specific objectives are as follows:

1. To assess the experience of key transition practices among young people with CP and their parent(s)/carer(s) in the Irish healthcare system.
2. To assess the provision of key transition practices to young people with CP, from the perspectives of disability service providers in the Irish healthcare system.
3. To explore the perspectives of young people with CP and their parent(s)/caregiver(s) on how to implement transition in the Irish healthcare system.
4. To explore the perspectives of service providers on transition for young people with CP.

**METHODS AND ANALYSIS**

We will use a convergent parallel mixed-methods design to address the overall aim of this study. Quantitative and qualitative data will be collected and integrated at the interpretation stage. This study will be conducted between January 2020 and December 2021. Figure 1 illustrates the planned timeline. Recruitment is ongoing.

**Framework of key transition practices**

The following framework of key transition practices was informed by a recent research programme conducted in the UK, the United Kingdom’s National Institute for Health and Care Excellence guideline on transition, and relevant Irish policy documents as practices that improve the experience and outcomes of transition from child to adult services (online supplemental appendix 1). The framework was also informed by discussions with members of the Project Management and Study Steering Groups, which includes health professionals, young people with CP and parents. This framework will be used to assess gaps in the current management of transition and to explore enablers, barriers and supports required to implement successful transition in the Irish healthcare system, for young people with CP.
1. A named worker, known to the young person, who oversees, coordinates or delivers transition support, acts as the link between the young person and the various practitioners involved in their care including their GP. This person may not be formally allocated and may not be a health provider but should be in close contact with health services.

2. Involvement of parent(s)/carer(s) in the young person’s care at a level that is deemed appropriate by both the young person and parent(s)/carer(s).

3. Child and adult services provide young people and families with information that describes the transition process and the support available before and after transfer. The information should specifically mention health services, which encompasses services that have a direct impact on people’s physical health, mental health and social well-being. This information should be provided early enough to allow young people time to reflect and discuss with parent(s)/carer(s) or health professionals and be in an accessible format. Where there is no adult service for a young person to transfer to, information about known and trusted voluntary organisations who could provide support should be provided to the young person.

4. Promotion of health self-efficacy (ie, actively helping young people to feel confident in managing their condition), where health encompasses complete physical, mental and social well-being, including provision of information to the young person about their condition and encouragement to take responsibility for their health.

5. Promotion of opportunities for self-management, where the individual is directly involved in planning and decision-making around their needs and takes responsibility for maintaining optimal physical, mental and social well-being.

6. A health professional from the relevant adult services or primary care meets the young person before they transfer from child services.

7. A senior manager with responsibility for implementing transition strategies and policies, including facilitating good working relationships between child and adult services, ensuring appropriate materials are available, and monitoring that the person has a suitable appointment in adult services. This person may not be known to the young person.

8. Where there is no adult service for a young person to transfer to, a detailed discharge letter is sent to the young person’s GP.

9. Formal training, relevant to health condition, in wider life skills—education, gaining employment, finances, housing, social relationships, sexual health, mental health. The health service may not provide such training but during consultations, staff should inquire about such matters and make referrals to other agencies as needed.

Patient and public involvement statement

A Study Steering Group of young people with CP, parent(s)/carer(s) and service providers will work closely with the Project Management Group and advise on all steps in the research processes. This will include developing study documentation such as information sheets, developing and piloting questionnaires, developing and piloting topic guides and interpreting findings. The Study Steering Group has informed the development of the framework of key transition practices. The group will also contribute to the development of materials to share findings with young people, families and service providers, such as an information leaflet and infographic. Further, based on the findings, the group will support the Project Management Group to identify and develop resources to facilitate successful transition for young people with CP in Ireland.

Participants

Participants will be young people with CP aged 16–22 years, their parent(s) or carer(s), and individuals who provide health services to young people with CP who are residing or working in Ireland. We will include people with CP of all severities of physical impairment.

Recruitment

Participants will be recruited from across Ireland via disability service providers, special education needs schools, universities, general practices, professional organisations and organisations for people with disabilities and through media, social media and word of mouth. Advertisements and participant information leaflets will include the URL to access an online version of the questionnaire and contact details for the research team, so that participants may request further information or the questionnaire in alternative formats.
Young people, their parents or carers and service providers who complete the questionnaire will be asked to contact a member of the research team if they would like to participate in an interview. We will develop a sampling frame and purposively sample young people, parents or carers and service providers for interview based on characteristics that may affect the experience or provision of transition. For young people, these may include age, gross motor function, intellectual disability and geographical location. For service providers, these may include the professional role of the individual, sector and geographical location.

**Quantitative data collection**

We will aim to collect quantitative data from up to 100 service users (ie, young people, parent(s), carer(s)) and service providers via a questionnaire. Participants may complete the questionnaire online, in person, by post or by telephone or video-conference. Young people may obtain support from a parent, family-member or carer to complete the questionnaire. To maximise inclusivity, we will make the questionnaire accessible to people with varying impairments including mild-to-moderate intellectual disability, for example by providing the survey in alternative formats and using various enhancements such as visual aids, simple words, clear questions and story technique. Where it is not possible to provide sufficient adaptations to obtain the young person’s perspectives, a parent or carer may complete the questionnaire on their behalf. The questionnaire was developed in collaboration with and piloted on young people, parents and service providers prior to use.

The following data will be collected from young people with CP:
1. Sociodemographic and condition-specific data.
2. Current and previous service use.
3. Experience of key transition practices.
4. Unmet health needs using an unmet needs questionnaire.\(^5\)

The following data will be collected from service providers:
1. Service-related data.
2. Predominant transition model.
3. Provision of key transition practices.

**Qualitative data collection**

Face-to-face, telephone or videoconference in-depth interviews will be conducted with service users and service providers. Approximately 25 service users and 20 service providers will participate in interviews. Service user interviews may be conducted individually with the young person or jointly with young people and their parent(s) or carer(s). If the young person is under 18 years, they must be interviewed with a parent or guardian present. Adaptations will be made within the interview to allow for inclusion of people with mild-to-moderate intellectual disability, communication impairment, visual impairment, hearing difficulty or other impairments. These may include providing them with the topic guide in lay language in advance, using alternative systems of communication and providing pictorial memory aids. Individual interviews with parent(s)/carer(s) will be facilitated where it is not possible to provide sufficient adaptations to obtain the young person’s perspectives. Interviews will be conducted at the participant’s home, workplace or other suitable and convenient venue.

We will use the framework of key transition practices to develop the topic guide for interviews. However, we will also explore participants’ perspectives of transition more broadly. Separate topic guides for service users and service providers will be developed in collaboration with and piloted on young people, parents and service providers.

Examples of topics to be explored during interviews with service users will include (1) experience of key transition practices; (2) if they believe these practices improved the experience of transition or in the case where practices were not experienced if these practices may have improved their experience; (3) how these practices should be implemented, for example, through structured programmes, written information, verbal information and (4) other supports that improved or would improve their experience of transition. Examples of topics to be explored during interviews with service providers will include (1) perceptions of successful and unsuccessful transition; (2) barriers and enablers to implementing key transition practices including organisational structures, processes and relationships; (3) readiness for implementation and (4) supports required to implement successful transition including the content and format of resources to facilitate implementation.

**Analysis**

Descriptive statistics will be used to report quantitative data such as sociodemographic data and experience of key transition practices. Where sufficient data are collected, we will examine associations between the experience of transition practices and sociodemographic and CP-related factors (eg, Gross Motor Function Classification System (GMFCS) level), respectively, using appropriate regression models. Associations between service provider characteristics and provision of key transition practices may also be explored using appropriate regression models where sufficient data are available.

Interviews will be analysed using the Framework Method, which allows for both deductive and inductive analysis of data. The Framework Method is appropriate for this study as we have predefined topics that we wish to explore but are also open to the emergence of additional themes. The Framework Method involves five iterative stages: familiarisation, thematic framework identification, labelling, charting and mapping and interpretation.\(^29\) A sample of transcripts will be read by two members of the research team until familiarity with the data is established. They will independently develop provisional codes before agreeing on an initial framework through discussion. The initial framework
developed will be applied to all transcripts and will be iteratively refined through our analytic process if new codes are identified. Analysis will be undertaken by one researcher. A proportion of coded transcripts will be checked by a second researcher. We will explore if demographic and CP-related factors (eg, GMFCS level) influence participants’ perspectives of transition. Data will be arranged into charts that summarise themes, issues, and individual responses. Finally, members of the Project Management Group will discuss emerging categories and themes collaboratively. Preliminary results and interpretations will also be shared with the study steering group who will have the opportunity to reflect and input on the findings. Strategies to enhance the trustworthiness of the findings, such as negative case analysis, peer-debriefing and reflexivity, will be used.

Integration
Integration of quantitative and qualitative data will be carried out at the interpretation stage of the research.30 We will use the key transition practices as a framework for developing themes and meta-themes. We will search for data related to each theme in questionnaires and interviews. We will also identify any additional emerging themes from questionnaires and interviews. We will use a convergence coding matrix to integrate themes and to develop meta-themes.31 This will involve grouping themes according to similar concepts and interpreting them to generate meta-themes. We will search for agreement and disagreement between studies according to meta-themes. Agreement and disagreement will be defined as convergence (ie, findings agree directly), complementarity (ie, findings offer complimentary information), dissonance (ie, findings seemingly contradict each other) or silence (ie, themes arise in quantitative data or qualitative data but not both). Integration will be supported by discussions with the Project Management Group and Study Steering Group.

Data management
A data management plan will be created prior to the start of data collection. All data will be stored securely. All participants will be assigned an ID number, which will be recorded on study documentation. Anonymous data entered electronically via Online Surveys will be downloaded securely into a study database. Data collected on paper forms will be entered into the study database and paper forms will be stored securely in a locked cabinet. Data entry and validation will be a continuous process. Audio files will be downloaded to an encrypted laptop on completion of interviews before being transferred to secure servers. Audio files will be transcribed verbatim by a professional transcription service and imported into qualitative data management software (Nvivo V.12: QSR International). Pseudonyms will be used to report participant quotes with consent. Identifying details will be removed to preserve participant anonymity.

Project management
The Project Management Group, consisting of the Chief Investigator (JR), co-applicants for the grant and post-doctoral researcher will be responsible for delivering all objectives.

ETHICS AND DISSEMINATION
This study has been approved by the RCSI University of Medicine and Health Sciences Research Ethics Committee (REC201911010). All participants will provide informed consent prior to data collection. For young people aged 16–18 years, a parent/guardian will also provide consent for the young person to participate. Examples of the participant information sheet (online supplemental material 1) and consent form (online supplemental material 2) are available in the online supplemental material. Information for potential participants is also shared on the study website https://www.ignitionstudy.com. We will use findings from this research, and discussion with the Study Steering Group, to identify resources that will facilitate successful transition from child to adult services for young people with CP in Ireland. A detailed dissemination plan will be developed in the early phases of the study. We will use a variety of knowledge translation activities to ensure that the outputs generated from this research are accessible to non-academic stakeholders including young people with CP, families and service providers. We will work closely with the Study Steering Group to identify and develop materials to share findings, such as an information leaflet and infographic summarising the findings and a seminar for young people, families and service providers. Results will be published in open access peer-reviewed journals and presented at national and international scientific conferences.

Author affiliations
1Department of Public Health and Epidemiology, RCSI University of Medicine and Health Sciences, Dublin, Ireland
2College of Health, Medicine and Life Sciences, Brunel University London, Uxbridge, UK
3School of Nursing and Midwifery, Queen’s University Belfast, Belfast, UK
4Medical Department, Central Remedial Clinic, Dublin, Ireland
5School of Nursing, Midwifery and Health Systems, University College Dublin, Dublin, UK
6Institute of Psychiatry, Psychology & Neuroscience, King’s College London, London, UK
7Physiotherapy Department, Central Remedial Clinic, Dublin, Ireland
8National Disability Children & Families Team, Social Care Division, Health Service Executive, Dublin, Ireland
9Office of the Chief Clinical Officer, Health Service Executive, Dublin, Ireland

Contributors All authors listed meet the International Committee of Medical Journal Editors criteria for authorship. JR and MW conceived the study, MN, OH, MO, TK, CK, AW, GL and MB designed the study. JR will lead the running of the study. JF will lead the collection, management and analysis of the data. MN and AW will lead the qualitative analysis. JR will lead the statistical analysis. All authors have read and approved the final manuscript.

Funding This work was supported by the Health Research Board, grant number APA-2019–004. The study sponsor is RCSI University of Medicine and Health Sciences.

Competing interests None declared.
Patient consent for publication  Not required.

Provenance and peer review  Not commissioned; externally peer reviewed.

Supplemental material  This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any error and/or omissions arising from translation and adaptation or otherwise.

Open access  This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.

ORCID iDs
Jennifer Fortune http://orcid.org/0000-0001-8971-1236
Aisling Walsh http://orcid.org/0000-0002-5312-5101

REFERENCES
17. NCEPOD. The national confidential enquiry into patient outcome and death, each and every need. London; 2018.