

Document Control

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V4.1	22/08/24	Draft	Version 4 updated with comments
V4.2	03/09/24	Draft	Transition meeting: add in Fontan forum, updated
			referral boxes, algorithm 1 – amended to say 1st
			visit in ACHD as required
V4.3	16/09/24	Draft	Agreed changes and signed off at SLT
V4.3	15/10/24	Draft	Sent to transition leads at AH and MFT. No
			transition Lead in post at MFT ah JR not in post at
			AH. Awaiting confirmation of who to approach
V4.3	19/12/24	FINAL	No feedback received from Transition team at AH
			and no one available at MFT – so SLT decision to
			proceed to ratification

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- > Laura Jole (ACHD Specialist Nurse MFT)



- > Psychology Team at Alder Hey Childrens Hospital NHS Foundation Trust
- Psychology Team at Liverpool Heart and Chest Hospital NHS Foundation Trust

Approved By: NWCHD Network Ratification Task and Finish Group's (Paediatric + ACHD) on behalf of the Network

Date: 20/01/2025

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Young People with Congenital Heart Disease

Protocol for transition and transfer of care between paediatric and adult congenital cardiology services in the North West of England, North Wales and The Isle of Man

Date: February 2025







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Executive Summary



This protocol has been developed to promote the planning and management of the care of young people with congenital heart disease during transition and transfer of care between paediatric and adult cardiology services. This protocol has been written under the auspices of the North West of England, North Wales and the Isle of Man CHD Network and meets the current NHS England standards (Section I) 2016.

Congenital-heart-disease-standards-and-specifications.pdf (england.nhs.uk)

1. Introduction

1.1 Purpose

- 1.1.1 Transition is defined in the Department of Health's 2006 publication *Transition: getting it right for young people* as "a purposeful, planned process that addresses the medical, psychosocial and educational/vocational needs of adolescents and young adults with chronic physical and medical conditions as they move from child centred to adult-oriented health care systems."
- 1.1.2 Congenital heart disease describes a range of heart conditions resulting from an abnormality of the heart structure or function that is present from birth. The majority of conditions are diagnosed in infancy or early childhood but occasionally it is not diagnosed until adulthood.
- 1.1.3 The aims of transition services are to ensure uninterrupted and coordinated care. To promote understanding of the patient's heart condition and its impact on adult life (higher education, employment, relationships, mental health and psychological wellbeing). To develop skills in self-care and promote independence and to support parents (DH, 2006).
- 1.1.4 This protocol has been developed to promote seamless transition services for young people with congenital heart disease who live in the North West of England, North Wales and the Isle of Man. This protocol is intended to:
 - Promote key standards and clear pathways of care from paediatric to adult congenital heart disease (ACHD) services.
 - Provide advice, guidance and support for the management of care during transition
 - Promote the appropriate transfer of care of young people to ACHD services
 - Build on existing relevant guidelines for routine care of young people focusing on areas where additional or different care should be offered with congenital heart disease
- 1.1.5 This protocol will help practitioners achieve a seamless transfer of care by providing a framework for working with young people with congenital heart disease at each of the major phase of adolescence. It is divided into four parts and provides:



- An overview of the service provision arrangements in the region, general principles and best practice
- A description of the key standards to consider when planning and delivering transition services
- The age and timings criteria for transition and transfer of care
- A practical framework to support the delivery of transitional care
- Patient questionnaires to support discussions with young people
- Patient Information Leaflet

1.2 Scope

- 1.2.1 This protocol applies both to paediatric and ACHD services.
- 1.2.2 This protocol applies to cardiology services for young people regardless of the complexity of their heart condition or where they live in the North West of England, North Wales or the Isle of Man.
- 1.2.3 This protocol is intended for healthcare staff involved in transition planning and delivery as well as the transfer of health care of young people with congenital heart disease.

1.3 Terminology

- 1.3.1 The term **'transition"** is used in this protocol to refer to the process of enabling a young person to become independent with their heart condition and to support their successful transfer of care to adult services. Transition has been divided into early, mid and late stages and should begin by the age of thirteen years. The term **"transition"** is not to be used in any patient facing publications following consultation with young people.
- 1.3.2 The term **'transfer'** or **'handover'** is used in this protocol to refer to an event within the process of transition when the care of the young person transfers from paediatric to adult health services.
- 1.3.3 The term "Young Person's Specialist Nurse" used in this protocol refers to a Specialist Congenital Heart Nurse who has the specific role of co-ordinating, managing and supporting young people through the transition process, transfer of care and until they have successfully been integrated into adult services. This service is nurse led and supports the young person throughout. They are referred to as YP Specialist Nurses to avoid using the term 'transition'.
- 1.3.4 The term **'young person/people'** is used in this protocol to refer to those between 13 years and 18+ years of age. It is widely recognised that it is good clinical practice to begin to discuss the need for transition early and that many young people will require support until the age of 25yrs.



- 1.3.5 The term "**Handover Clinic**" is used in this protocol to refer to the formal handover clinic run jointly with the paediatric and ACHD service. This facilitates formal hand over of care to adult services for all complex patients and may include patients with moderate CHD where there is a concern that intervention or surgery may be required in the near future.
- 1.3.6 The term "Young Person's Clinic" or "YP Clinic" is used in this protocol to refer to the clinics run by the YP Specialist Nurses to support the young person throughout the transition process from age 13yrs to 18yrs+ regardless of how their care is transferred. This may be in the form of one-to-one clinic sessions, or they may be held as group sessions held either in the specialist centres or locally throughout the region. Every young person should be offered at least one appointment in the Young Person's clinic recognising that some patients will require several appointments during their transition journey.
- 1.3.7 The term "ACHD" is used to refer to Adult Congenital Heart Disease services.
- 1.3.8 The term "**learning disability**" is used throughout this document and in line with national policy is a formal diagnostic term for a person who has global and pervasive difficulties with social and adaptive functioning, present before the age of 18. A learning disability affects the way a person understands information and how they communicate. This means they can have difficulty understanding new or complex information, learning new skills and coping independently. A learning difficulty, which is different to a learning disability, would encapsulate a specific difficulty in one domain, e.g. dyslexia.
- 1.3.9 The term "Neurodiversity" will be used to describe neurodivergent conditions including Autism/Autism Spectrum Condition (ASC), Autism Spectrum Disorder (ASD) and Attention Deficit Hyperactivity Disorder (ADHD).

1.4 Current service provision arrangements

- 1.4.1 Young people (up to the age 18 years) growing up with a congenital heart problem are cared for by the Specialist Paediatric Cardiology Teams based at Alder Hey Children's Hospital NHS Foundation Trust and Royal Manchester Children's Hospital, Manchester University NHS Foundation Trust. These two specialist centres also support 26 peripheral clinics throughout the North West of England, North Wales, and the Isle of Man so that young people can be seen nearer to home.
- 1.4.2 From year 6 all patients with a single ventricle and Fontan circulation are invited to a **"Fontan Forum"** to support their transition from primary school to high school. This is a special event run by the YP specialist nurses and psychologists at Alder Hey Children's Hospital and is available to all patients across the Network.
- 1.4.3 All young people with congenital heart disease will be supported by a named YP Specialist Nurse throughout the process and will be invited to attend a "Young Person's Clinic" at either
 - Alder Hey Children's NHS Foundation Trust



- Royal Manchester Children's Hospital (Manchester University NHS Foundation Trust)
- 1.4.4 Young people with complex congenital heart disease or patients with moderate disease that is unstable will be invited to a **"Handover Clinic"**. These clinics will be held at either
 - Alder Hey Children's NHS Foundation Trust
 - Royal Manchester Children's Hospital (Manchester University NHS Foundation Trust)
- 1.4.5 Young people with simple and moderate congenital heart disease that is stable will have their care transferred from the Consultant Paediatric Cardiologist to a Consultant ACHD Cardiologist based at one of the following hospitals:
 - Liverpool Heart and Chest NHS Foundation Trust (Level 1 centre)
 - Manchester Heart Centre, Manchester University NHS Foundation Trust (Level 2 centre)
 - Wythenshawe Hospital, Manchester University NHS Foundation Trust (Level 3)
 - Blackpool Victoria Hospital, Blackpool, Fylde and Wyre Hospitals NHS Foundation Trust (Level 3 centre)
 - Wrexham Maelor Hospital, Betsi Cadwaladr University Health Board, North Wales (Level 3 centre)

1.5 General principles

- 1.5.1 Care of young people during transition and the transfer of their care between paediatric and adult congenital cardiology services should be planned and managed. Paediatric and adult congenital cardiology healthcare teams should work jointly during transfer of care.
- 1.5.2 Management of young people with congenital heart disease should consider their needs and preferences.
- 1.5.3 Communication between healthcare professionals (in paediatric and adult services) and young people and their family is essential. It should be supported by transition assessments tailored to the young person's needs.
- 1.5.4 General practice has a crucial role to play as this will remain consistent throughout the transition and transfer of care to adults. A new enhanced service for general practice was introduced in 2014/15 to ensure proactive care and personalised care planning for people with complex health and care needs who may be at high risk of unplanned admission to hospital. This will be under the supervision of a named, accountable GP. All documentation related to transition should be copied to the patients GP.
- 1.5.5 Where patients have complex needs involving several specialist teams and professionals involved. The YP Specialist Nurse will help co-ordinate be-spoke MDT meetings as



- appropriate to ensure good communication amongst the teams involved. This is to ensure that transitional care is coordinated and planned.
- 1.5.6 If emergency care is required for their congenital heart problem during the transfer stage the paediatric CHD team would still be responsible for providing care up until the time the patient has been seen in adult services for the first time. This needs to be effectively communicated with the patient and family and documented on the patient's medical notes and transition plan. There may be individual circumstances that require emergency care to be provided by the adult service. Where this decision has been made it is imperative that this is clearly documented and shared. Once the patient has been seen in adult services all future emergency admissions would be managed within adult services. It is important that the patient and family understand how to manage urgent care during this transfer stage and beyond.
- 1.5.7. NHSE CHD Standards (2016) state that referral should occur from age 12yrs. However NICE Guidance (2016 and updated in 2023) suggests that referral to transitional services should occur in year 9 at school (between the age of 13-14yrs). The NW CHD Network has therefore adopted the NICE Guidance in writing this protocol.
- 1.5.8 Referrals to Transitional services at Alder Hey NHS Foundation Trust and Royal Manchester Children's Hospital, Manchester NHS Foundation Trust should be made by copying in the Specialist Nurses into relevant clinic letters from the age of 13 years. Patients seen in peripheral clinics should be referred to the appropriate transition service linked to either Alder Hey or Royal Manchester Children's Hospital.

Young Person's Clinic at RMCH

Referral letter and/or copy of clinic letter to
be sent to
Maria Hollingsworth
Dr Giovanni Ciotti
Royal Manchester Children's Hospital
Oxford Road
Manchester M13 9WL

Young Person's Clinic at Alder Hey

Referral letter and/or copy of clinic letter to be sent to Amanda Haworth Dr Rob Johnson Alder Hey Children's Hospital East Prescot Road Liverpool L14 5AB

1.6 Key standards for implementation

NHSE Congenital Heart Disease Standards and Specifications (2016)

1.6.1 NHS England published and ratified a set of service standards for transition and resource requirements for the designation of specialist services for patients with congenital heart disease. The standards related to transition are detailed below:

NHSE Transition Standards (Section I)

Congenital Heart Networks must demonstrate arrangements to minimise loss of patients to follow-up during transition and transfer. The transition to adult services will



be tailored to reflect individual circumstances, taking into account any special needs. 'Lost to follow-up' rates must be recorded and discussed at the network multidisciplinary team meeting.

- All services that comprise the local Congenital Heart Network must have appropriate arrangements in place to ensure a seamless pathway of care, led jointly by paediatric and adult congenital cardiologists. There must be access to beds and other facilities for adolescents.
- There will not be a fixed age of transition from children's to adult services but the process of transition must be initiated no later than 12 years of age, taking into account individual circumstances and special needs.
- All patients requiring long-term congenital care undergoing transition must be seen at least once for consultation by an ACHD cardiologist and an ACHD Specialist Nurse, in a specialist multidisciplinary team transfer clinic or equivalent. Clear care plans/transition passports must be agreed for future management in a clearly specified setting, unless the patient's care plan indicates that they do not require long-term follow-up.
- Patients, partners, families and carers must be fully involved and supported in discussions around the clinical issues in accordance with the patient's wishes. The views, opinions and feelings of the patient must be fully heard and considered, and the patient must be offered the opportunity to discuss matters in private, away from their parents/carers if they wish.
- All patients transferring between services will be accompanied by high quality information, including the transfer of medical records, imaging results and the care plan.
- Young people undergoing transition must be supported by age-appropriate information and lifestyle advice. Management of young people arriving in the adult service will aim to ensure that they are fully confident in managing their own condition and health care. In the clinic, they will see an ACHD Specialist Nurse who will explain and discuss a range of issues including the impact of their condition, contraception and pregnancy, and lifestyle, in language the young person can understand. The Cardiologist will discuss the treatment plan with the young person and discuss it with their family/carers when appropriate. The young person will have some independent time to talk with their Specialist ACHD Cardiologist and ACHD Specialist Nurse.
- **18** The particular needs of young people with learning disabilities and their parents/carers must be considered, and reflected in an individual tailored transition plan
- Young people must have the opportunity to be seen by a Practitioner Psychologist on their own. Psychological support must also be offered to partners/family or carers.



2. Overview of the protocol

Wherever possible the protocol is in the format of algorithms and frameworks, supported by minimum text. The component parts of the transition and transfer of care pathway for young people with congenital heart disease are discussed in three management stages as follows and the overall pathway is outlined in **Algorithm 1**:

3. Early, mid, and late-stage transition

The early stage describes the period from when the decision to prepare for transition is made, usually within paediatric care. The mid phase is that of transition readiness when the young person and their parents/carers are supported to become more independent and knowledgeable. The late stage occurs when the young person not only transfers their care to adult services but is actively participating in adult care activities e.g. self-management and decision making. (Telfair etc 1l, 2004).

Table 1

OVERVIEW OF THE PROTOCOL

Early-Stage Transition

This transition pathway identifies young people with congenital heart disease from 13 years to 14 years.

Mid-Stage Transition

This transition pathway identifies young people with congenital heart disease from 15 years to 16 years.

Late-Stage Transition

This transition pathway identifies young people with congenital heart disease from 17 years to 18 years who will require transfer from paediatric to adult cardiology services

ALGORITHM 1 (OVERVIEW)

TRANSITION AND TRANSFER OF CARE PATHWAY (AGE 13 TO 18 YEARS)

From age 13yrs



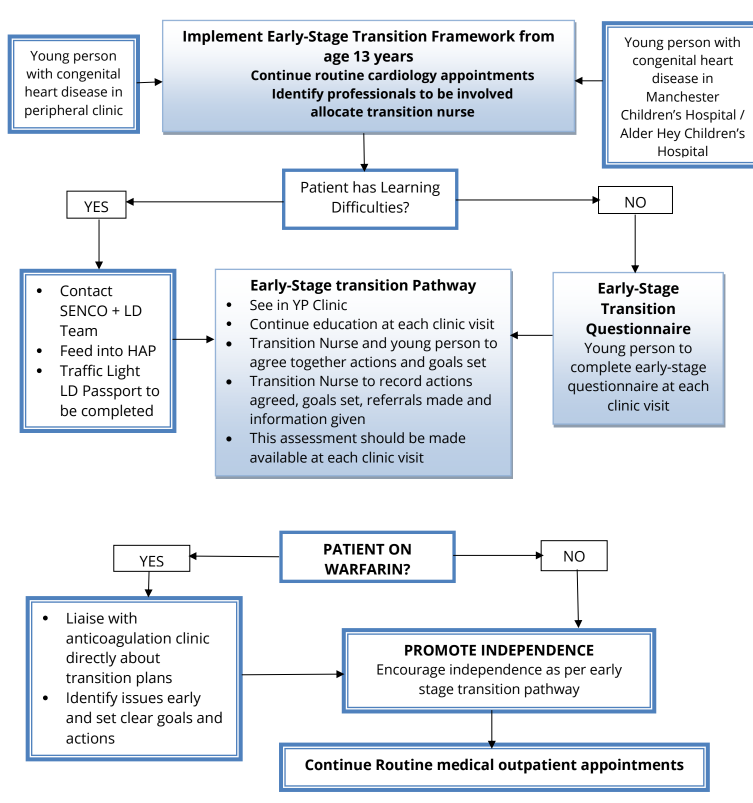
At age 13yrs Simple, Moderate & Complex Congenital Heart Disease • Discuss transition and give leaflet **Manchester Children's Hospital** Alder Hey Children's Hospital **Peripheral Clinics in DGHs** • Refer to Transition service + invite to Young Person's Clinic Continue routine medical OPA **Transitional Care** Young Persons' Congenital Cardiac Clinic Age 13yrs to 18yrs Simple, Moderate & Complex Congenital Heart Disease Nurse Led Clinics: **Manchester Children's Hospital** Alder Hey Children's Hospital • Continue pathway and when ready care to be transferred Introduce handheld notes Simple ACHD **Moderate ACHD Complex ACHD** Transition Clinic Transfer of care - direct **Transition Clinic Cardiology Clinic** Age 16yrs to 18yrs Age 16yrs to 18yrs **Complex ACHD and unstable moderate ACHD** Simple and moderate stable ACHD Paediatric Cardiology & Adult Congenital Heart Teams **Paediatric Cardiologist to ACHD Royal Manchester Children's Hospital Cardiologist** Alder Hey Children's Hospital • Meet ACHD Team and continue pathway Meet ACHD Team and continue pathway Commence management plan Commence management plan **Transfer of Care** (cc all correspondence to ACHD Nurse Specialists Invited to visit ACHD service prior to first appointment. YP nurse specialist to attend 1st appt if required (pathway completed)

Paediatric Cardiology Clinics

EARLY-STAGE TRANSITION: 13 YEARS UP TO 14 YEARS



ALGORITHM 2



EARLY-STAGE TRANSITION: 13 YEARS UP TO 14 YEARS



Recommended management

Algorithm 2 describes the recommended pathway for young people with congenital heart disease aged 13 years up to 14 years of age.

- 4.1 The paediatric cardiac nurse should start to introduce the concept of preparing to transfer care to adult health care with the young person and their family by age 13 years. They should given a copy of the patient information leaflet which briefly describes the transition process (NWCHDN_11.1 "Moving On"). This should be documented in the patients' notes, with a plan to formally commence the transition process at age 13 years.
 - Moving On to Adult Services Northwest, North Wales, Isle of Man Congenital Heart Network (northwestchdnetwork.nhs.uk)
- 4.2 The early-stage transition pathway should start by the age of 13 years. The young person should be offered an appointment in the nurse led Young Persons Congenital Cardiac Clinic.
- 4.3 The aims of early-stage transition can be facilitated by using a planning framework which identifies the roles and responsibilities of those involved in the process (see table 2). From the start of the transition process, all organisations have a responsibility to work together to ensure that information is shared to inform each other's assessments and that coherent planning is based on a real understanding of the young person's needs.
- 4.4 The transition assessment/plan provides a checklist for discussion with the young person at each of the three stages of transition: early (Doc: NWCHDN_1.3.1), middle (Doc: NWCHDN_11.3.2) and late (Doc: NWCHDN_11.3.3). The checklists for each key area suggest some general aims and subjects for discussion, although additional areas may be identified to meet individual young people's needs.
- 4.5 The young person's level of understanding should be assessed by using advanced communication skills and open questioning techniques.
- 4.6 At early-stage transition, the young person should become aware of their own heart condition and health and care needs, lifestyle issues including how they are coping at school etc. An assessment of the young persons' level of understanding is important.
- 4.7 The patient questionnaire for early-stage transition can be used in the Young Person's Clinic to support this process: <a href="https://www.nwchan.edu.ne.gov/nwchan.edu.n
- 4.8 Any actions agreed, goals set, referrals made, or information (such as leaflets) given to the young person during the discussion should be recorded.
- 4.9 This should be completed by the YP Specialist Nurse and agreed with the young person and reviewed at least annually thereafter. This assessment should be made available



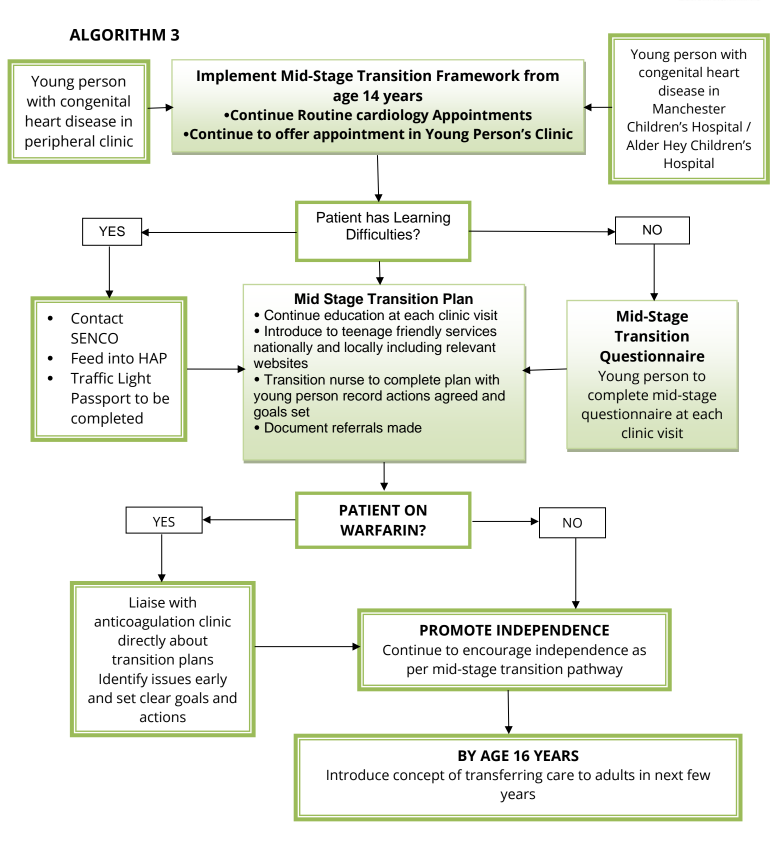
- for all clinic appointments throughout early transition. Once completed, the early-stage transition assessment should be filed in the young person's medical notes.
- 4.10 The concept of seeing a professional on their own should be gradually introduced to the young person to give the family time to adjust to this change. However, it must be emphasised that the young person can *choose* who is in the clinic appointment with them. Parents must always remain involved and should be seen with the young person at some time during the session.

Planning Framework for early-stage transition (Table 2):

A Planning Framework for Early-Stage Transition From age 13 years		
What Should Happen?	Who Should Do It?	
Outpatients – all young people age 13 years to be	Medical staff (including Paediatric	
referred to nurse led transition service	Cardiologist or SpR)	
Copy transition team into correspondence	Clinic Nurses	
In-patients - all young people age 13 years to be	Paediatric Cardiac Liaison Nurse	
referred to nurse led transition service	(PCLN)	
	Ward based nurses	
Give leaflet "Moving On" to patient and discuss	Referrer	
with patient what is involved (Doc: NWCHDN_11.1)		
Patient invited to attend Young Persons' Congenital	YP Specialist Nurse	
Cardiac Clinic		
Begin "early-stage" transition pathway		
Give out early-stage transition questionnaire at		
each clinic visit		
Set up handheld notes and give to patient at first		
appointment		
Support family and young person through this		
early stage		
Contact any other relevant members of the MDT		
and ensure that their contact details are		
documented in medical notes		
Ensure all members of team are aware of referral including GP		

MID STAGE TRANSITION: AGE 14 YEARS UP TO 16 YEARS







Recommended management

Algorithm 3 describes the recommended pathway for young people with congenital heart disease aged 15 years up to 16 years of age.

- 5.1 The middle-stage transition pathway should be started by the age of 14 years.
- 5.2 The aim of middle-stage transition can be facilitated by using a planning framework which identifies the roles and responsibilities of those involved in the process (Table 3).
- 5.3 Discussion with the young person should be recorded in their medical notes.
- 5.4 During mid-stage transition the young person and their family should be given information about medical, educational and psychosocial development to enable the young person to set goals for participating in their own care.
- 5.5 Young people should be helped to take responsibility for their own medications from as early an age as possible. This stage of the process will also prepare and equip the family for the growing independence and eventual transfer of care to adult services.
- 5.6 The young person's level of understanding should be assessed by using advanced communication skills and open questioning techniques.
- 5.7 The patient questionnaire for middle stage transition should be used in the Young Person's Clinic to support this process NWCHDN 11.4.2 Mid-Stage-Questionnaire FINAL 02-08-21.docx (live.com)
- 5.8 Any actions agreed, goals set, referrals made, or information (such as leaflets) given to the young person during the discussion should be recorded.
- 5.9 This should be completed by the YP Specialist Nurse and the young person and reviewed at least annually thereafter. This information should be made available for all clinic appointments throughout middle stage transition and be recorded in the patients medical notes.
- 5.10 The concept of transfer to adult cardiology services should be introduced by 15 years at the latest to adequately prepare the young person and their parents for this.
- 5.11 It is acknowledged that some patients who have special needs will have a statement of educational need and will already be going through a formal transition process from the age of 14 years. This will involve their educational, developmental, and social care needs, but will also address any health-related issues. It is envisaged that this pathway will sit comfortably alongside any such documentation and compliment any multiagency work that is on-going. This will ensure that their congenital heart needs are fully met.



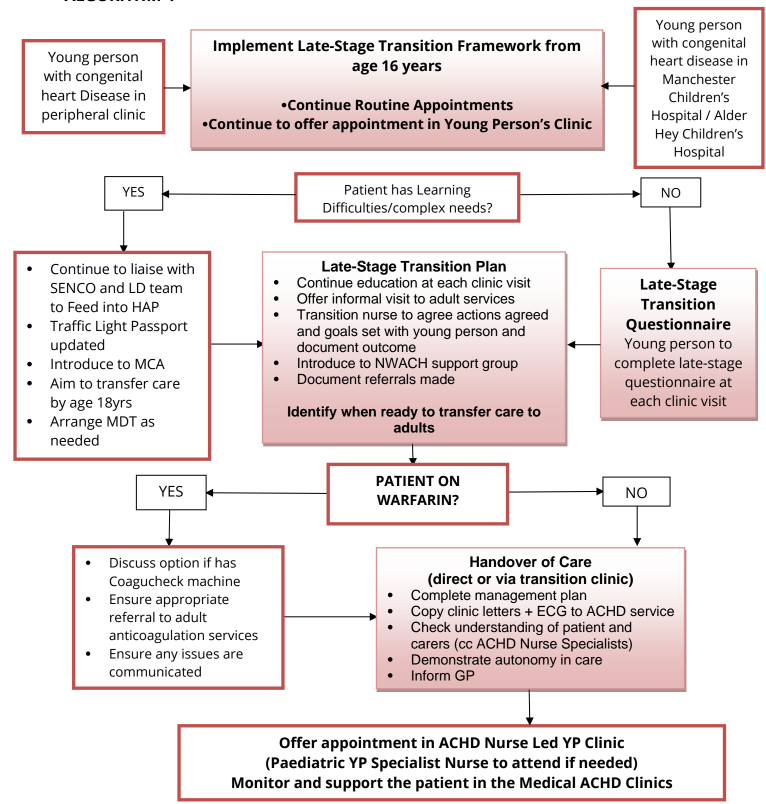
Planning framework for mid-stage transition (table 3)

A Planning Framework for Mi	d-Stage Transition
What Should Happen?	Who Should Do It?
Routine medical appointments to continue as	Medical staff
normal	
Offer appointments in Young Person's clinic as	YP Specialist Nurse
appropriate	
Give out mid stage transition questionnaire at the	
beginning of every appointment	
Continue with mid-stage transition pathway	
Introduce to teenage friendly services including	
BHF, LHM and TSHF	
If on warfarin – liaise with anticoagulation service	YP Specialist Nurse
and GP for review of how they are managing and	
issues with compliance	
If patient has learning difficulties and has an	YP Specialist Nurse/SENCO
educational statement, contact SENCO at school to	Educational staff
ensure that health issues are included in their	
transition process	
Be involved in the development of Health Action	YP Specialist Nurse/SENCO
Plans for those young people with learning	
difficulties	
If learning difficulties - ensure patient has a traffic	YP Specialist Nurse
light passport document in case of in-patient	
admissions	
Ensure appropriate referrals made to MDT as	YP Specialist Nurse
appropriate and as highlighted during clinic	
assessments	

LATE-STAGE TRANSITION: AGE 16 YEARS AND UP TO AGE 18 YEARS+



ALGORITHM 4





LATE-STAGE TRANSITION AND TRANSFER OF CARE: AGE 16 YEARS AND UP TO AGE 18 YEARS+

Recommended management

Algorithm 4 describes the recommended pathway for young people with congenital heart disease aged 16 years and up to 18 years+.

- 6.1 Late-stage transition should be started by the age of 16 years.
- 6.2 The aims of late-stage transition can be facilitated by using a planning framework which identifies the roles and responsibilities of those involved in the process (Table 4).
- 6.3 Discussion with the young person at late-stage transition should be recorded using the transition assessment/plan (see **Doc: NWCHDN_11.3.3**).
- 6.4 The young person and their family should be feeling confident about leaving paediatric services and the young person should have a considerable degree of autonomy over their own care. The young person's level of understanding should be documented.
- 6.5 Any actions agreed, goals set, referrals made or information (such as leaflets) given to the young person should be recorded.
- The patient questionnaire for late-stage transition can be used to support this process.

 NWCHDN 11.4.3 Late-Stage-Questionnaire FINAL 02-08-21.docx (live.com)
- 6.7 The transition assessment/plan should be completed by the YP Specialist Nurse and the young person. Both the YP Specialist Nurse and the young person should sign the late-stage transition record. This assessment should be recorded in the patient's medical records.
- 6.8 This assessment will follow the patient into adult services.

7. Transfer of care to adult services

- 7.1 All young people including those with learning disabilities or special needs should be supported to leave paediatric services by the age of 18 years where appropriate.
- 7.2 A referral to the formal Handover Clinic will be required in the following situations:
 - a. Patients with complex CHD
 - b. patients where it is anticipated that they may require surgery or intervention in the next 12 months
 - c. patients with learning disabilities and complex needs
 - d. Patients with complex needs and multiple specialities involved may need a be-spoke MDT meeting to be arranged.



- 7.3 In the Handover Clinic a formal medical handover of care will be facilitated through an MDT approach with both Paediatric and ACHD Cardiologists in attendance. The YP Specialist Nurse and ACHD Nurse Specialist will also be available to support the patient and family.
- 7.4 A detailed clinic letter is required for all young people transferring care to adult services regardless of how their care is transferred. This should be copied to the ACHD Nurse Specialists and must include as a minimum:

Minimum information required at handover (Table 4)

Information required	Details
Demographics	Include parents/carers details State if has LD or special needs
Cardiac Diagnosis	e.g. Fallot – RVEDv abs/indexed + PR + EF +/- stenotic elements coarctation – presence of stenosis/aneurysm AR root dimensions etc
Co-morbidities to include developmental, neuro-psychological/cognitive and mental health issues	Include contact details of all other services involved in care
Summary of Surgical Procedures	Include operation notes
Summary of cardiac catheter procedures	Include reports
Specialist Investigations (MRI/CT) dates and key points	Include relevant reports
List of current medications and recent changes prior to transfer	
Current status at time of transfer	

- 7.5 A comprehensive management plan will be given to the young person and a copy sent to their GP.
- 7.6 For young people with moderate and simple ACHD a referral should be made directly to the routine ACHD Clinic. A detailed clinic letter will be sent, and a copy of the young person's management plan will be given to the young person and a copy sent to their GP. They will be supported throughout the process by the YP Specialist Nurse in the same way.



Table 5

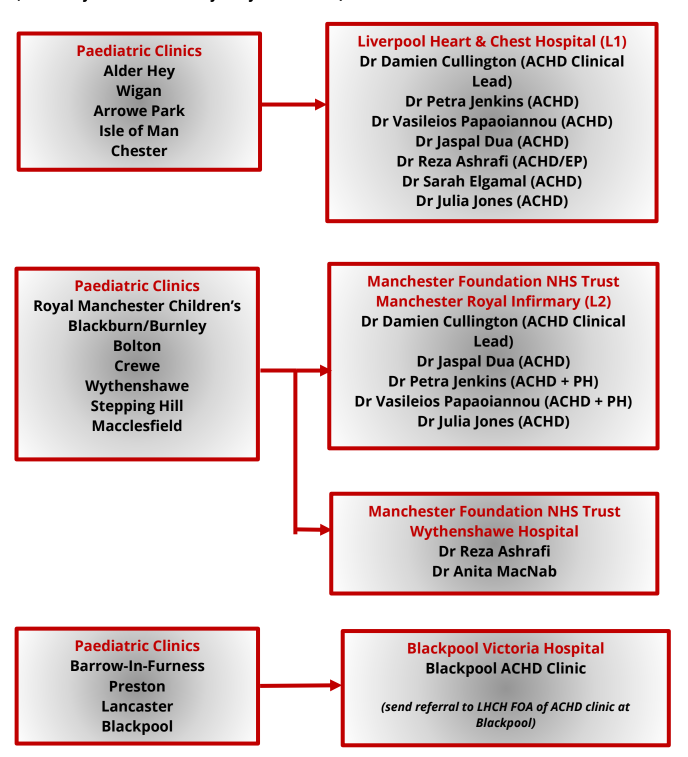
A Planning Framework for Late-Stage	TI di i Sicioni
What Should Happen?	Who Should Do It?
Routine medical appointments to continue as normal	Medical staff
Offer appointment in Young Person's Clinic as appropriate	YP Specialist Nurse
Give out late-stage transition questionnaire at the beginning of	
every appointment	
Continue with late-stage transition pathway	
Offer informal visit to adult services	
If on warfarin – liaise with anticoagulation service for review of	YP Specialist Nurse
how they are managing and issues with compliance and identify	
where follow-up management will occur	
If patient has a learning disability or neurodiversity and has an	YP Specialist Nurse/SENCO/
educational statement continue to work with SENCO at school to	Educational staff
ensure that health issues are included in their transition process	
If a patient has a learning disability or neurodiversity – make sure	YP Specialist Nurse
that they understand the implications of the Mental Capacity Act	
once they are 16yrs of age	
Be involved in the development of Health Action Plans for those	YP Specialist Nurse/SENCO
young people with learning difficulties	
If learning difficulties - ensure patient has a traffic light passport	YP Specialist Nurse
document in case of in-patient admissions	
Ensure appropriate referrals made to MDT as appropriate and as	
highlighted during clinic assessments. If required coordinate an	
MDT meeting with all relevant teams in attendance to co-	
ordinate care and timings of transfer	
When ready ensure patient is either referred to Handover Clinic	
for formal handover of care or liaise with their Paediatric	
Cardiologist to inform them of their readiness for direct transfer	
of care	
Ensure ACHD management plan completed and given to patient	YP Specialist Nurse/ACHD
	nurse specialist
Ensure ACHD patient information leaflets are given to patient	YP Specialist Nurse
	ACHD nurse specialist
Give copies of latest ECG and clinic letters to patient	YP Specialist Nurse
	Consultant Paediatric
	Cardiologist
Ensure patient understands decisions made and the plan for	Paediatric and Adult
future care. This is to include plans for emergency care during	Congenital Cardiologist
the transfer of care.	and YP Specialist Nurse/
	ACHD Nurse Specialist
Support and help parents/carers through this process	YP Specialist Nurse/
	ACHD Nurse Specialist
Ensure referral letter sent to appropriate Adult Cardiologist as	Paediatric Cardiologist/
per agreed plan and copied to GP	Paediatric Cardiac Secretary
Ensure all relevant documentation is transferred to appropriate	Paediatric Cardiac secretary
adult services	



Ensure new appointment issued in adult service	ACHD team secretary
YP Nurse Specialist to follow care through to adults for at least	YP Specialist Nurse
first appointment if required	
Sign off transition pathway once completed	ACHD Nurse Specialist

8. Guide to aid referral to adult services

(Please refer to APPENDIX A for referral details)





Paediatric Clinics Bangor Wrexham ACHD clinic Dr Raj Thaman Glan Clywd Abersywyth Wrexham Maelor Hospital Wrexham ACHD clinic Dr Petra Jenkins (LHCH)

Patients to be referred directly to ACHD Clinics at Level 1 (L1) and Level 2 (L2)

Patients with complex CHD should be referred centrally to MRI or LHCH this includes

- Fontan
- Palliated CHD
- Cyanotic patients
- Pulmonary hypertension
- Pts where upcoming surgery is planned
- Pts with learning disabilities or significant comorbidities

Inherited Cardiac Conditions and Aortopathy

Patients with Inherited Cardiac Conditions (ICC) should be referred to the regional ICC clinics at Liverpool Heart and Chest Hospital or Manchester Royal Infirmary or adult cardiologist looking after other family members. Refer to NWCHDN_52_ICC transition SOP

Patients with aortopathy can be referred to ICC or ACHD

Cardiac Maternity Clinics

Joint Obstetric Cardiac Clinics (JOCC)

St Mary's Hospital Manchester

Dr Anna Roberts/Prof Bernard Clarke/Dr Damien Cullington/Prof Bernard Keaveney

Liverpool Women's Hospital

Dr Naomi McGuiness/Dr Vishal Sharma/Dr Damien Cullington/Dr Reza Ashrafi

9. Auditing and monitoring

- 9.1 Activity levels in the following clinics needs to be recorded and reported to the NW CHD Network
 - The Young Person's Clinic



- The formal Handover Clinic
- ACHD Nurse Led Transition Clinic
- ACHD Medical Clinic
- 9.2 Quality Indicators will be monitored locally and nationally via a national patient satisfaction survey
- 9.3 Monitoring needs to include loss to follow up rate.



Appendix A

ACHD Clinics and Contact Details

(please ensure ACHD Nurse Specialist are copied into correspondence)

Hospital	Contact	Telephone Number
Liverpool Heart & Chest Hospital NHS Foundation Trust Thomas Drive Liverpool L14 3PE (ACHD Level 1 Centre for North West of England, North Wales and Isle of Man)	ACHD Cardiologists: Dr Damien Cullington (Clinical Lead) Dr Sarah Elgamal (imaging) Dr Jaspal Dua Dr Petra Jenkins Dr Reza Ashrafi (EP) Dr Vasileios Papaioannou Dr Julia Jones	Switchboard: 0151 600 1616
	Anna Harrison and Fiona Andrews Specialist Nurse - Adult Congenital Heart Disease	Tel: 0151 254 3333 Mobile: 07542229889
Manchester University Foundation Trust, Manchester Heart Centre, MRI, Oxford Road, Manchester, M13 9WL (ACHD Level 2 Centre for North West of England, North Wales and Isle of Man)	ACHD Cardiologists Dr Damien Cullington (Clinical Lead), Dr Jaspal Dua Dr Petra Jenkins (PH) Dr Vasileios Papaioannou (PH and general ACHD) Dr Julia Jones Cardiologists with a special interest in ACHD Professor Bernard Clark	Switchboard: 0161 276 1234
	Laura Jole, Melanie Menzies, and Kate Shammas Specialist Nurses - Adult Congenital Heart Disease	0161 276 1234 Vocera: ask for ACHD Nurse
Manchester University Foundation Trust, University Hospital of South Manchester Southmoor Road, Wythenshawe M23 9LT	ACHD Cardiologist Dr Reza Ashrafi Cardiologist with a Special Interest in ACHD Dr Anita MacNab	Switchboard: 0161 998 7070



		NORTH MEST NORTH WALLS
(ACHD Level 2 Centre for North West of England, North Wales and Isle of Man)	Laura Jole, Melanie Menzies, and Kate Shammas Specialist Nurses - Adult Congenital Heart Disease	0161 276 1234 Vocera: ask for ACHD Nurse
Blackpool Fylde and Wyre Hospital, Whinney Heys Road, Blackpool,	ACHD Cardiologist (Refer to LHCH FAO of Blackpool ACHD)	Tel: 0151 254 3350
FY3 8NR, (ACHD Level 3 Centre for Lancashire and Cumbria)	Specialist Nursing Team – Adult Congenital Heart Disease via the helpline at Liverpool Heart & Chest Hospital	Tel: 0151 254 3333
Betws Cadwalader University Health Board – Wrexham Maelor Hospital, Croesnewydd Road, Wrexham, LL13 7TD (ACHD Level 3 Centre for North	Consultant Cardiologist with an Interest in ACHD Dr Raj Thaman ACHD Cardiologist Dr Petra Jenkins (Based at LHCH)	Tel: 01978 727 801
Wales)	Nicola Coates Specialist Nurse - Adult Congenital Heart Disease	Tel: 01978 727 801



Appendix B

Comments from young people and their families on current and future transition services in the region

Prior to the development of this protocol a workshop was held in May 2005 with young people and their families to discuss current and future transition services. This workshop highlighted particular issues that have been considered in the development of this protocol.

This is what young people and their families said about the current transition process:

- The transition process is not always understood by young people, their parents/carers or professionals, and the referral pathways are not always clear.
- Young people and their families do not have one person they can call to assist them through the transition.
- The transition process can feel very stressful because there appears to be a lack of planning ahead.
- The expectations of young people are often not well managed, and they are not always well informed.

This is what young people and families said would improve the transition process:

- The transition plan should be more holistic and include social, employment, leisure and education opportunities.
- Young people want to feel that they are listened to as an equal partner in the transition planning.
- Young people and their parents/carers want to have access to a key worker who oversees the whole process and guides them through the transition.
- Young people want to have consistent, accessible and easy to understand information on the transition process and on their condition, medication, treatments and tests to help them make their own choices.
- The young person's voice should always be listened to and be at the centre of the process.



Appendix C

Details of Formal Transition Clinics at Royal Manchester Children's Hospital and Alder Hey NHS Foundation Trust

Frequency of clinics will increase as the CHD service expands

Paediatric (cardiology and adult congenital cardiology transition clinics are held for young people lex ACHD at 16 years to 18 years to support transfer of care.
Who	Paediatric Cardiologist, Adult Congenital Cardiologist, Paediatric Cardiac Nurse Specialist/Transition Nurse and Adult Congenital Heart Nurse Specialist and cardiac physiologists. Cardiology Clinic staff
Where	Main outpatient's department within Manchester Children's Hospital NHS Foundation Trust
When	Formal handover (Transition) clinic once a week Young People's clinics once a week
How	All referral letters to be copied to the Transition Nurse Specialist for co-ordination of care

Alder Hey	Children's NHS Foundation Trust	
Paediatric cardiology and adult congenital cardiology transition clinics are held for young people		
with compl	ex ACHD at 16 years to 18 years to support transfer of care.	
Who	Paediatric Cardiologist, Adult Congenital Cardiologist, Paediatric Cardiac Nurse Specialist/Transition Nurse, Adult Congenital Heart Nurse Specialist and cardiac physiologist. Cardiology clinic staff	
Where	Main outpatient department within Alder Hey Children's NHS Foundation Trust	
When	Formal handover (Transition) clinic once a week	
	Young People's clinics 3 x a week	
How	All referral letters to be copied to the Transition Nurse Specialist for co-ordination of care	



Appendix D

List of Abbreviations

ACHD Adult Congenital Heart Disease

DH Department of Health

ECG Electrocardiogram – a test that records the heart's electrical activity

Echo Echocardiogram – a test that looks at the structure and function of the heart

FE Further Education

GP General Practitioner

GUCH Grown Up Congenital Heart Disease

INR International Normalized Ratio – a blood test which measures the time it takes

for blood to clot and compares it to an average

LD Passport A learning disability hospital passport A hospital passport provides important

information about a patient with a learning disability, including personal details, the type of medication they are taking, and any pre-existing health conditions. The passport also includes information about how a person communicates and their likes and dislikes. Health Guides: Hospital Passports, Summary Care Records And

Flu Jabs | Mencap

MDT Multi-Disciplinary Team

N/A Not Applicable

NHS National Health Service

NICE National Institute Clinical Excellence

NSF National Service Framework

PCLN Paediatric Cardiac Liaison Nurse

PE Physical Education

POC Point of Care – a term used for a test which is performed for or by a patient

outside of the usual laboratory setting

SENCO Special Educational Needs Coordinator

SpR Specialist Registrar



Appendix E

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