Public health functions to be exercised by NHS England

Service specification No.18
NHS Sickle Cell and Thalassaemia Screening Programme
This specification is part of an agreement made under the section 7A of the National Health Service Act 2006. It sets out requirements for an evidence underpinning a service to be commissioned by NHS England for 2014-15. It may be updated in accordance with this agreement.
Public health functions to be exercised by NHS England

Service specification No.18
NHS Sickle Cell and Thalassaemia Screening Programme

Prepared by –
Public Health England
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Service specification No.18

This is a service specification within Part C of the agreement ‘Public health functions to be exercised by NHS England’ dated November 2013 (the ‘2014-15 agreement’).

The 2014-15 agreement is made between the Secretary of State for Health and NHS England under section 7A of the National Health Service Act 2006 (‘the 2006 Act’) as amended by the Health and Social Care Act 2012.

This service specification is to be applied by NHS England in accordance with the 2014-15 agreement. An update to this service specification may take effect as a variation made under section 7A of the 2006 Act. Guidance agreed under paragraph A38 of the 2014-15 agreement may inform the application of the provisions of this service specification.

This service specification is not intended to replicate, duplicate or supersede any other legislative provisions that may apply.

The 2014-15 agreement including all service specifications within Part C is available at www.gov.uk (search for ‘commissioning public health’).
Section 1: Purpose of Screening Programme

1.1 Purpose of the Specification

To ensure a consistent and equitable approach across England a common national service specification must be used to govern the provision and monitoring of the linked antenatal and newborn NHS Sickle Cell and Thalassaemia (SCT) Screening Programme.

The purpose of the service specification is to outline the service and quality indicators expected by NHS England (NHS E) for NHS England responsible population and which meets the policies, recommendations and standards of the UK National Screening Committee (UK NSC).

The service specification is not designed to replicate, duplicate or supersede any relevant legislative provisions which may apply, e.g. the Health and Social Care Act 2008 or the work undertaken by the Care Quality Commission. The specification will be reviewed and amended in line with any new guidance as quickly as possible.

This specification needs to be read in conjunction with the following:

- SCT Screening Programme Standards [http://sct.screening.nhs.uk/standards](http://sct.screening.nhs.uk/standards)
- NHS Newborn Blood Spot screening programme service specification
- Specialised Haemoglobinopathy Services definition


- Guidance and updates on KPIs [http://www.screening.nhs.uk/kpi](http://www.screening.nhs.uk/kpi)
- UK NSC guidance, Managing Serious Incidents in the English NHS National Screening Programmes [http://www.screening.nhs.uk/quality-assurance#fileid9902](http://www.screening.nhs.uk/quality-assurance#fileid9902)
1.2 Screening programme aims, objectives and health outcomes

The SCT Programme aims to:
Support people to make informed choices during pregnancy and before conception
Improve infant health through prompt identification of affected babies
Ensure high quality, accessible care throughout England
Promote greater understanding and awareness of the conditions and the value of screening

1.2.1 Antenatal Sickle Cell and Thalassaemia Screening Programme

Aim:
To offer timely antenatal sickle cell and thalassaemia screening to all women (and couples), to facilitate informed decision-making.

Outcome to be achieved – for those women accepting prenatal diagnosis, 50% of prenatal diagnoses to be performed before 12 weeks 6 days.

1.2.2 Newborn Sickle Cell Screening Programme

Aim:
To identify babies born with conditions where early intervention is likely to be beneficial

Outcome - to achieve the lowest possible childhood death rate and to minimise childhood morbidity from sickle cell diseases.

1.2.3 Linked Antenatal and Newborn Sickle Cell and Thalassaemia Screening Programme

Aim: to link results from antenatal tests taken by parents-to-be with their baby’s test result

1.3 Objectives

- To ensure an appropriate level of understanding about screening and these conditions among professionals involved with the programme.
• To minimise the adverse effects of screening.

1.4 Public Health Outcomes Framework

SCT screening contributes to the Public Health Outcomes Framework indicator on the uptake of screening for national screening programmes. Indicator 2.2iii Access to non-cancer screening programmes: Sickle cell and thalassaemia screening

1.5 Principles

All individuals will be treated with courtesy, respect and an understanding of their needs,

All those participating in the sickle cell and thalassaemia screening programme will have adequate information on the benefits and risks to allow an informed decision to be made before participating.

The target population will have equitable access to screening.

Screening will be effectively integrated across a pathway including between the different providers, screening centres, primary care and secondary care.
Section 2: Scope of Screening Programme

2.1 Description of screening programme

The UK NSC policy on antenatal screening for sickle cell disease and thalassaemia is that all eligible women should be offered screening.

The SCT Programme is comprised of two, linked, screening programmes: sickle cell and thalassaemia screening during pregnancy; and sickle cell screening offered to all newborns in England as part of the NHS Newborn Blood Spot Screening Programme.

2.2 Care pathway

A description of both the antenatal SCT screening pathway and the newborn sickle cell element of the NHS Newborn Blood Spot Screening pathway are given below, along with diagrams of the pathways showing failsafe processes identified by the national screening programme. The Provider is expected to follow the pathway.


Antenatal Sickle Cell and Thalassaemia Screening Programme

The pathway for the antenatal screening for Sickle cell and Thalassaemia as part of the linked antenatal and newborn screening programme consists of the following:

- The eligible population is identified through routine midwifery-led antenatal care or primary care and offered screening by 10 weeks.
• Midwives provide written information and take consent.

• All women (in both high prevalence (HP) and low prevalence (LP) Trusts) are offered a screening blood test for thalassaemia using the initial results from routine red blood cell indices.

  - In high prevalence areas, all women are also offered a screening blood test for sickle cell and other haemoglobin variants.

  - In low prevalence areas, the Family Origin Questionnaire (FOQ) is used as an initial screening tool to identify any woman and the baby’s father who are from a high-risk group for haemoglobin variants. If the questionnaire shows that there is a risk of either parent being a carrier, a screening blood test is offered to the woman.

• Pathways to be in place to ensure women who are known carriers (or carrier couples) can be referred directly to appropriate counselling service immediately without being delayed by routine antenatal booking processes to ensure that PND, if wanted, can happen very early in pregnancy (this may involve additional provider contracts or referral to genetic counseling).

• Laboratory tests the sample as per national policy and reports results preferably using antenatal status codes. This can result in one of four outcomes:
  - Mother is identified as a carrier: healthcare professional informs parents of results and screening is offered to the baby’s father. Results should be included on the newborn blood spot card for ongoing pregnancies.
  - Nothing abnormal is detected on screening: mother’s results are included on the blood spot card.
  - Inconclusive (or incomplete) parental results: screening is offered to the baby’s father
  - Affected mother (sickle cell disease): mother is referred to a consultant for clinical and obstetric management and screening is offered to the baby’s father.

• If both parents are confirmed carriers and/or either is affected they are referred as an at-risk couple for pre-natal diagnostic testing and offered counselling by trained counsellors, see (sct.screening.nhs.uk/externalraining).

• If the baby’s father is not available for testing, prenatal diagnosis can be offered if wished on the basis of the mother’s carrier status.

• Parents who decline pre-natal diagnostic testing continue with the pregnancy. Results should be included on blood spot card for ongoing pregnancies.

• If prenatal diagnosis testing is accepted, parental samples are taken and sent to the specialist laboratory for testing.
- If PND result is normal or the fetus is identified as a carrier and the pregnancy continues, the PND result should be recorded on the blood spot card.
- If PND testing identifies that the fetus is affected with a major haemoglobin disorder, the parents are provided with information relating to the specific disorder. Informed choice about continuation of pregnancy is offered. If they choose to continue with the pregnancy, the results are recorded on the blood spot card. If they choose not to continue with the pregnancy rapid access to termination of pregnancy services is required.

- For pregnancies resulting in a live birth, the antenatal screening pathway ends when the antenatal screening results are included on the blood spot card (screening and PND results)
- For women opting for termination of pregnancy, the antenatal screening pathway ends when the women are counselled appropriately following a termination of pregnancy
Public health functions to be exercised by NHS England

Antenatal Sickle Cell and Thalassaemia Screening, Including Failsafes

Failsafe SCT 2
Offer and documentation

Failsafe SCT 3
All eligible women who accept screening are screened

Failsafe SCT 7
Receipt of screening sample by the laboratory

Failsafe SCT 9/14
Inconclusive or incomplete parental results

Failsafe SCT 10/17
Reporting antenatal carrier results

Failsafe SCT 12
Offer of screening to baby’s father

Nothing abnormal detected on screening

Inconclusive (or incomplete) parental results

Carrier result

Affected mother (sickle cell disease) – refer to consultant for clinical & obstetric management

Baby’s father available and gives consent

Baby’s father not available or declines consent (with conclusive maternal result)

Laboratory tests sample and reports results as per local arrangements

Nothing abnormal detected on screening of baby’s father

Confirmed carrier or affected result in both parents – refer at risk couple
Newborn Sickle Cell Screening Programme within Newborn Blood Spot Screening

The pathway for newborn screening for sickle cell is an integral part of the NHS Newborn Blood Spot Screening Programme and consists of the following:

- The eligible population is identified through NN4B issued at birth or registration with a GP practice for babies born abroad.
- Midwives check antenatal results and family history. Ideally all antenatal results obtained from antenatal SCT screening are included on the blood spot card.
- Midwives provide written information (ideally before birth) and take consent.
- Screening can be offered to unscreened babies who move into a local area up to one year of age. The Health Visitor is responsible for offering screening to parents of babies with no written evidence of screening results. The child health record departments who note the arrival of a baby (when it is registered) alert the HV to unscreened babies.
- Samples are taken routinely on day 5 and in exceptional circumstances between day 5-8, (day of birth is day 0), in accordance with Guidelines for Newborn Blood Spot Sampling, and sent to the appropriate newborn screening laboratory. Records are kept of all tests including those declined.
- A pre transfusion sample to screen for sickle cell disease is taken on all babies admitted to a neonatal unit. The blood spot card should be marked “pre transfusion”.
- The “pre transfusion” blood spot card should be stored with the baby’s medical record in line with local protocols and dispatched to the newborn screening laboratory together with the routine 5 day sample if the baby has received a blood transfusion in the interim.
- As a failsafe, transfused babies who did not have a pre transfusion sample taken before transfusion can be tested for sickle cell disease using DNA analysis. Such samples are sent by the newborn screening laboratory to one of two DNA laboratories. This service is commissioned nationally until October 2014, when responsibility transfers to NHS England.
- Newborn screening laboratory tests sample as per national policy and reports all results to Child Health Records Department (CHRD).
- Screen positive results are also reported to local clinician/and designated sickle cell and thalassaemia centre (under development). The designated sickle cell and thalassaemia centre ensure that affected babies enter the care pathway (refer to The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services 2011), and return diagnostic results back to the newborn screening laboratories to confirm enrollment into care.
Laboratory testing results in one of four outcomes:

- Condition not suspected: parents are informed of the result

- Baby is identified as a carrier: results are reviewed against maternal (and paternal) results where these are available (to assist communication and identify any cases where misdiagnosis or non-paternity could be an issue) then parents are informed ideally by face-to-face discussion, or by letter with offer of a face-to-face session

- Inconclusive result: additional sample may be required

- Condition suspected: immediate clinical referral initiated and parents informed of the result. The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services (2011) specifies a failsafe system to ensure all screen positive babies enter the care pathway

- CHRDs maintain a list of the eligible population to provide a failsafe check to identify untested babies by 17 days, to monitor coverage and to send results to health visitors and parents according to national policy.

See Section 3.13 for details of the end of the screening pathway.
Public health functions to be exercised by NHS England

Newborn Sickle Cell Programme within Newborn Blood Spot Screening, Including Failsafes

Failsafe NBS 4
Identifying the eligible population

Failsafe NBS 5
Offer

Failsafe NBS 17a
Coverage

Failsafe NBS 17b
Identification of untested babies

Failsafe NBS 21
All babies with screen positive results are offered diagnostic test

Failsafe NBS 22
Communicating newborn screening results to parents
2.3 Failsafe arrangements

Quality Assurance (QA) within the screening pathway is managed by including failsafe processes. Failsafe is a back-up mechanism, in addition to usual care, which ensures if something goes wrong in the screening pathway, processes are in place to (i) identify what is going wrong and (ii) what action follows to ensure a safe outcome.

In accordance with UK NSC standards and protocols the provider is expected to:

- have appropriate failsafe mechanisms in place across the whole screening pathway. A complete list of the failsafe processes in the SCT Screening Programme to be met by the Provider can be found on the national SCT screening programme website and the Newborn Blood Spot Programme Website
- review and risk assess local screening pathways in the light of national SCT Screening Programme guidance
- work with NHS England and Quality Assurance Teams to develop, implement, and maintain appropriate risk reduction measures
- ensure that mechanisms are in place to regularly audit implementation of risk reduction measures and report incidents
- ensure that appropriate links are made with internal governance arrangements, such as risk registers
- ensure routine staff training and development
  - follow guidance from The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services to ensure all screen positive babies enter the care pathway

2.4 Roles and accountability through the screening pathway

The linked SCT screening programme is dependent on systematic specified relationships between stakeholders. Stakeholders include maternity units, the antenatal, newborn and pre-diagnosis screening laboratories, diagnostics laboratory and genetics services, child health records departments, and specialist sickle cell and thalassaemia services, i.e. ‘the screening pathway’. NHS England will be responsible for ensuring that the pathway is robust. For their part the provider will be expected to fully contribute to ensuring that cross-organisational systems are in place to maintain the quality of the whole screening pathway. This will include, but is not limited to:

- provision of robust coordinated screening that ensures all parties are clear of their roles and responsibilities, so that there is clarity of handover of responsibility throughout all elements of the screening pathway
• ensuring that community midwifery services are supported to facilitate early booking for maternity care within primary and community care settings
• agreeing and documenting roles and responsibilities relating to all elements of the screening pathway across organisations including CHRDs
• developing joint audit and monitoring processes
• agreeing joint failsafe mechanisms where required to ensure safe and timely processes across the whole screening pathway
• contributing to any of NHS England and public health screening lead initiatives in screening pathway development in line with UK NSC expectations
• providing or seeking to provide robust electronic links with relevant organisations
• links with primary, secondary and tertiary care
• the need for robust IT systems across the screening pathway
• joint review meetings across the screening pathway to be held on a regular basis
2.5 Commissioning arrangements

The commissioning of the SCT screening pathway will involve commissioning different elements of the pathway at different levels.

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Provider</th>
<th>Level of commissioning</th>
<th>Level of contracting</th>
<th>Rationale and other comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANTENATAL</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Identify cohort in a timely manner</td>
<td>Maternity services / primary care</td>
<td>AT</td>
<td>CCG</td>
<td>The eligible population is identified through routine midwifery-led antenatal care</td>
</tr>
</tbody>
</table>
| Inform the cohort and maximize uptake of screening in a timely manner | Maternity services / primary care | AT                     | CCG                  | Informing the cohort and maximising uptake in a timely manner takes place during routine midwifery-led antenatal care and also sometimes in primary care.  
**CCGs will have responsibility for commissioning maternity care.** |
<p>| Screening test - sample taking / | Maternity services / primary care | AT                     | CCG                  | Mostly carried out through routine midwifery                                                   |</p>
<table>
<thead>
<tr>
<th>Pathway</th>
<th>Provider</th>
<th>Level of commissioning</th>
<th>Level of contracting</th>
<th>Rationale and other comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family Origin Questionnaire (FOQ) completed (antenatal)</td>
<td></td>
<td></td>
<td></td>
<td>care.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Midwives take the blood sample and ensure the FOQ is completed and send to labs. GPs may also take the blood sample.</td>
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<td></td>
<td></td>
<td>CCGs will have responsibility for commissioning maternity care.</td>
</tr>
<tr>
<td>Screening test - analysis</td>
<td>Antenatal labs</td>
<td>AT</td>
<td>CCG</td>
<td>Level of commissioning needs to be above that of individual CCGs to allow centralisation of these services and to improve quality and efficiency as per direction of the pathology modernization agenda. (Carter Review of NHS Pathology Services (2006, 2008) made the case for consolidating pathology nationally to improve quality, patient safety and</td>
</tr>
<tr>
<td>Pathway</td>
<td>Provider</td>
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<td>Level of contracting</td>
<td>Rationale and other comments</td>
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<tr>
<td>Results reporting</td>
<td>Maternity services or specialist counselling service</td>
<td>AT</td>
<td>CCG</td>
<td>Results reporting as part of routine midwifery care. Negative blood test results given to mother/couple by midwife. Positive blood test results given to mother/couple by specialist counsellor/midwife. In some areas, especially high prevalence areas, specialist haemoglobinopathy counsellors carry this out. Giving of positive results includes advice about testing the baby's father. Maternity services to be commissioned by CCGs</td>
</tr>
<tr>
<td>Counselling of “at-risk” couples</td>
<td>Maternity services/Specialist Nurses/Genetic Centre</td>
<td>AT and Linked with specialised commissioning</td>
<td>NHS England/CCG</td>
<td>Counselling carried out by specialist midwives, haemoglobinopathy counsellors, genetic</td>
</tr>
<tr>
<td>Pathway</td>
<td>Provider</td>
<td>Level of commissioning</td>
<td>Level of contracting</td>
<td>Rationale and other comments</td>
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<tr>
<td></td>
<td>Specialist haemoglobinopathy DNA centres (currently three in England)</td>
<td>Specialist haemoglobinopathy DNA centres (currently three in England)</td>
<td>Specialist haemoglobinopathy DNA centres (currently three in England)</td>
<td>counsellors and occasionally obstetricians who also need to be appropriately trained. Arrangements need to include couples identified from previous pregnancies as being “at risk”, who would be directly referred for counselling at start of pregnancy and not go through the usual ‘booking process’. Two of the three specialist genetic centres provide counselling and cvs/amniocentesis for local services as well as DNA analysis of specimens.</td>
</tr>
<tr>
<td>Sample taking - amnio/CVS sample taking</td>
<td>Fetal medicine specialists Specialist haemoglobinopathy DNA centres</td>
<td>Fetal medicine specialists Specialist haemoglobinopathy DNA centres</td>
<td>Fetal medicine specialists Specialist haemoglobinopathy DNA centres</td>
<td>CVS is in the Specialised Services National Definition Set SSNDS. Amniocentesis where the procedure is</td>
</tr>
<tr>
<td>Pathway</td>
<td>Provider</td>
<td>Level of commissioning</td>
<td>Level of contracting</td>
<td>Rationale and other comments</td>
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<tr>
<td>(currently three in England)</td>
<td>(currently three in England)</td>
<td></td>
<td></td>
<td>difficult/complex is in the SSNDS Two of the three specialist genetic centres provide amnio/CVS sample taking. NHS England will have responsibility for commissioning specialised services (consistent with FA and Downs etc)</td>
</tr>
<tr>
<td>Sample analysis – Prenatal Diagnosis (PND)</td>
<td>Specialist genetic centres</td>
<td>NHS England – link to specialised commissioning at hub level</td>
<td>NHS England</td>
<td>This analysis is only carried out at three specialist genetic centres (Oxford, UCH and Kings) and involves small numbers (approx 400 samples p/a). Consider commissioning through one hub to ensure consistency.</td>
</tr>
<tr>
<td>Results reporting and counselling for women undergoing</td>
<td>A range of providers – all maternity services should</td>
<td>AT</td>
<td>CCG</td>
<td>Mostly carried out by specialist haemoglobinopathy</td>
</tr>
<tr>
<td>Pathway</td>
<td>Provider</td>
<td>Level of commissioning</td>
<td>Level of contracting</td>
<td>Rationale and other comments</td>
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<tr>
<td>PND</td>
<td>specify how this works for couples who have had a prenatal diagnosis either specialist haemoglobinopathy counsellor, specialist centre (where PND is done there) or obstetrician/specially trained midwife/genetic counselor</td>
<td></td>
<td></td>
<td>counsellor in high prevalence areas, but can be by other healthcare professionals.</td>
</tr>
<tr>
<td>NEWBORN</td>
<td></td>
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</tr>
</tbody>
</table>
| Newborn – identify cohort     | Maternity services                                                                 | AT                     | CCG                  | The eligible population is identified through the issuing of NN4B at birth (or registration with a GP practice for babies born abroad).  
  *CCGs will have responsibility for commissioning maternity care.*                                                                                                                                                                 |
<table>
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<th>Rationale and other comments</th>
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</thead>
<tbody>
<tr>
<td>Inform the cohort and maximize uptake of screening</td>
<td>Maternity services</td>
<td>AT</td>
<td>CCG</td>
<td>Informing the cohort and maximizing uptake takes place during routine midwifery-led antenatal and postnatal care, and sometimes through primary care. Health visitors inform families moving into the area.</td>
</tr>
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<td></td>
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<td></td>
<td>CCGs will have responsibility for commissioning maternity care.</td>
</tr>
<tr>
<td>Sample taking (part of Newborn Blood Spot Screening Programme)</td>
<td>Maternity services</td>
<td>AT</td>
<td>CCG</td>
<td>Midwife takes blood sample for sickle cell disease as part of Newborn Blood Spot screening. Neonatal/paediatric care services are responsible for taking the sample if a</td>
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<tr>
<td>Pathway</td>
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<td>------------------------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>Screening test – analysis (for sickle cell)</td>
<td>Newborn labs</td>
<td>specialised commissioning or Lead Area Office Commissioner for each Lab</td>
<td>NHS England</td>
<td>Link with specialised services for 13 newborn labs (and specialist haematology laboratories carrying out second line testing specifically for sickle cell)</td>
</tr>
<tr>
<td>DNA analysis (sickle cell) for transfused babies</td>
<td>DNA analysis reference labs for transfused babies (currently not in the bloodspot specification)</td>
<td>NHS England</td>
<td>NHS England</td>
<td>Two labs carry out this work (Sheffield and Kings) funded nationally until 2013. Funding needs to be picked up from 2013. Proposal has been to public health leads of specialist commissioners group and 'support/agreement in principle’ of need – chair</td>
</tr>
<tr>
<td>Pathway</td>
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<td>Level of contracting</td>
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<td></td>
<td>SCG chair from London.</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>[Two laboratories carry out this work (Sheffield and Kings) funded nationally until September 2014. Funding needs to be picked up from October 2014. Proposal has been to public health leads of specialist commissioners group and ‘support/agreement in principle’ of need – chair SCG chair from London. The SCT Programme has funded this for 2 years but funding will finish in 2014. This should now be commissioned as an integrated part of the</td>
</tr>
<tr>
<td>Pathway</td>
<td>Provider</td>
<td>Level of commissioning</td>
<td>Level of contracting</td>
<td>Rationale and other comments</td>
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</tr>
<tr>
<td>Results reporting</td>
<td>Child Health</td>
<td>NHS England</td>
<td>NHS England</td>
<td>mainstream newborn screening pathway for sickle cell disease from April 2014.</td>
</tr>
<tr>
<td></td>
<td>Records</td>
<td></td>
<td></td>
<td>Screen negative results: health visitor gives results to parents</td>
</tr>
<tr>
<td></td>
<td>Health visitors</td>
<td>NHS England</td>
<td></td>
<td>Carrier results: specially trained linked health visitor or trained counsellor give SCT carrier results to parents</td>
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<td></td>
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<td></td>
<td>Positive results: contact by specialist counsellors (usually in high prevalence areas or local clinical service (low prevalence areas)</td>
</tr>
</tbody>
</table>
The commissioning of the SCT screening pathway involves commissioning at different levels. The SCT screening services will be commissioned by NHS England alongside specialised services where appropriate.
2.6 Links between screening programme and national programme centre expertise

Public Health England (PHE) will be responsible for delivery of the essential elements of screening programmes best done once at national level.

These include:

- developing, piloting and roll-out to agreed national service specifications of all extensions to existing screening programmes and new screening programmes;
- setting and reviewing programme standards;
- setting and reviewing national service specifications and advising on section 7A agreements (under the direction of DH requirements);
- developing education and training strategies;
- providing parent information;
- determining data sets and management of data, for example to facilitate KPIs are collected;
- setting clear specifications for equipment, IT and data;
- procurement of equipment and IT where appropriate;
- (procurement may be undertaken by NHS England but will need advice from PHE screening expertise and related clinical experts);
- collect, collate and quality assure data for cancer and non-cancer screening programmes;
- monitor and analyse implementation of NHS commissioned screening services;
- provide advice to DH on priorities and outcomes for NHS England mandate and section 7a agreement, and to lead on detailed provisions, in particular the 7a agreement on screening;
- advise NHS England how to increase uptake of screening.

PHE will also be responsible for:

- providing the quality assurance functions for screening programmes;
- providing Public Health expertise and advice on screening at all levels of the system, including specialist Public Health expertise being available as part of NHS England screening commissioning teams;
- ensuring action is taken to optimise access to screening programmes, e.g. among socio-economically disadvantaged groups.
• Ensuring reports on important aspects of screening are available at various geographies (e.g. local authority, maternity service) to enable population based oversight
Section 3: Delivery of Screening Programme

3.1 Service model summary

Model summary

The linked national screening programme consists of:

- Antenatal screening offered to pregnant women early in pregnancy to identify women and then couples who are at increased risk (1:4) of an affected pregnancy to offer them the choice of prenatal diagnosis and the option of termination of an affected pregnancy or continuation of the pregnancy. This should ideally all take place before 12 weeks 6 days of pregnancy. Women who already know their carrier status (for example from a previous pregnancy) should be offered direct and speedy referral to counsellors for assessment of the couple risk status and prenatal diagnosis.

- All women (in both high prevalence (HP) and low prevalence (LP) Trusts) are offered a screening blood test for thalassaemia using the initial results from routine red blood cell indices.

- In high prevalence areas, all women are also offered a screening blood test for sickle cell and other haemoglobin variants.

- In low prevalence areas, the Family Origin Questionnaire (FOQ) is used as an initial screening tool to identify any woman and the baby’s father who are from a high-risk group for haemoglobin variants. If the questionnaire shows that there is a risk of either parent being a carrier, a screening blood test is offered to the woman.

- Newborn screening offered for sickle cell disease to all newborn babies as one of five conditions now tested for on the newborn blood spot (heel prick).
  - A pre-transfusion sample should be offered to babies at risk of having a blood transfusion. Transfused babies who do not have a pre transfusion sample taken can be tested for sickle cell disease using DNA analysis.
  - Additional tests are offered if required by, screening protocol to achieve a conclusive result.
  - Parents may decline all or part of the test.
  - A national IT failsafe solution should be in place to ensure samples are received in the laboratory and no babies born in England miss
being offered screening. To be effective this needs central commissioning.

- Conclusive results are recorded on a child health information system for the eligible population and for all five conditions. There needs to be a systematic notification of results to parents and the screening results recorded in the PCHR.
- All screen positive babies should enter into appropriate care which includes access to a designated clinician and relevant health professionals who confirm diagnosis and initiate appropriate clinical management and treatment. See The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services.
- All parents of babies with carrier results should be informed, ideally face to face, of their babies result.

- As with all newborn and antenatal screening the process of the offer of screening is largely embedded within the routine maternity and newborn pathway and not as a separate service.

There are key points about this programme which make it different, and also are relevant to effective commissioning. The key points for the SCT programme are:

- the importance of early testing in pregnancy to enable women to exercise choice as well as the possibility that testing may have already been done.
- recognition of the impact of lifetime genetic information. As an increasing proportion of women and their partners are aware of their carrier status before pregnancy, the choice of direct access for PND rather than routine pregnancy care should be available.
- the interface between maternity, laboratories, specialist counselling service and specialist diagnostic services
- the importance of timely and reliable communication by newborn screening laboratories of screen positive results to the local clinician/and the designated sickle cell and thalassaemia centre to ensure that affected babies enter the clinical care pathway. See The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services
- linkage with primary care and CHRDs.

All elements of the screening pathway should be delivered by appropriate staff and to national standards and guidelines.
Public health functions to be exercised by NHS England

3.2 Service model summary

In accordance with UK NSC standards and protocols the provider will be responsible for ensuring that the part of the programme they deliver is coordinated and interfaces seamlessly with other parts of the programme with which they collaborate, in relation to timeliness and data sharing.

The Provider will provide one or more named individuals who will be responsible for the coordination of the delivery of the programme and provider contribution to planning supported by appropriate administrative support to ensure timely reporting and response to requests for information. Where there is only one named coordinator, the provider will ensure that there are adequate cover arrangements in place to ensure sustainability and consistency of programme.

The Provider and NHS England will meet at regular intervals (at least annually). The meetings will include representatives from programme coordination, clinical services, laboratory services and service management.

3.3 Clinical and corporate governance

In accordance with UK NSC standards and protocols the provider will:

- ensure co-operation with and representation on the local screening oversight arrangements/structures
- ensure that responsibility for the screening programme lies at Director-level, (or delegated responsibility)
- ensure that there is appropriate internal clinical oversight of the programme and have its own management and internal governance of the services provided with the appointment of a Clinical Lead, a Programme Manager and the establishment of a multidisciplinary steering group (that meets quarterly) as a minimum
- ensure that there is regular monitoring and audit of the screening programme, and that, as part of organisation’s Clinical Governance arrangements, the organisation’s Board is assured of the quality and integrity of the screening programme
- comply with the UK NSC guidance on managing serious incidents
- have appropriate and timely arrangements in place for referral into treatment services that meet programme standards
- be able to provide documented evidence of clinical governance and effectiveness arrangements on request
- ensure that an annual report of screening services is produced which is signed off by the organisation’s Board
- have a sound governance framework in place covering the following areas:
3.4 Definition, identification and invitation of cohort/eligibility

The target population to be offered screening antenatally is all pregnant women, and the fathers of babies whose mothers are carriers or affected.

Women do not need to be tested again in the same or subsequent pregnancy provided that:

- There are two or more previous results which are from a reputable laboratory, preferably accredited by a body in the UK, which are consistent, unequivocal and well documented. These results must be interpreted in line with recommendations in the third edition of the laboratory handbook.

And

- The red cell indices remain the same

And

- The patient identification has three or more matching data items, e.g.
  - name, date of birth and NHS number/hospital number
  - name, date of birth and address
  - NHS number/hospital number, date of birth and address (if woman confirms name change)
  - name, date of birth and haemoglobinopathy card

For more information about the criteria please refer to the Third edition of the laboratory handbook.

The decision about re-testing should be made by the laboratory who will always need to perform routine blood indices. Requests and blood samples should be taken and sent as usual to the laboratory. Known carrier women should be referred directly to counselling services without waiting for checking of the routine indices.

The target population to be offered sickle cell screening as part of the NHS Newborn Blood Spot Screening Programme is all babies between 5 and 8 days of age and babies moving in to the country up to one year of age.

The Provider will make every effort to maximise screening uptake from vulnerable and hard-to-reach groups within the eligible population. This includes babies born abroad who move into the country up to one year of age.
3.5 Location(s) of programme delivery

The location of the offer of screening by midwives is to be locally determined.

Antenatal laboratory tests to be provided by laboratories meeting required laboratory standards which can be found on the National Screening programme website.

Pre-natal diagnostic testing is to be provided by specialist centres.

3.6 Days/Hours of operation

To be locally determined. However, timeliness is essential and is a key criterion of quality along all parts of the screening pathway.

3.7 Entry into screening programme

Antenatal: through GPs or direct referral into Maternity Services. While there is nothing specific in the GP contract regarding the SCT Programme, general practitioners have a key role in ensuring that pregnant women referred to them are referred on as soon as possible to Midwifery Services and for holding results of newborn screening.

Newborn: through midwifery services or through GPs/CHRDs for babies born abroad.

3.8 Working across interfaces

The screening programme is dependent on strong working relationships (both formal and informal) between professionals and organisations along the screening pathway. Accurate and timely communication and handover across these interfaces is essential to reduce the potential for errors and ensure a seamless pathway for service users. It is essential that there remains clear named clinical responsibility at all times and at handover of care the clinical responsibility is clarified. NHS England will be responsible for ensuring that the pathway is robust. For their part the Provider will ensure that appropriate systems are in place to support an interagency approach to the quality of the interface between these services. This will include, but is not limited to:
• agreeing and documenting roles and responsibilities relating to all elements of the screening pathway across organisations
• providing strong clinical and managerial leadership and clear lines of accountability
• developing joint audit and monitoring processes
• working to nationally agreed Programme standards and policies
• agreeing jointly on what failsafe mechanisms are required to ensure safe and timely processes across the whole screening pathway
• contributing to any NHS ENGLAND Screening Lead’s initiatives in screening pathway development in line with UK NSC expectations
• meeting the SCT screening programme standards covering managing interfaces which can be found in *NHS Sickle Cell and Thalassaemia Screening Programme: Standards for the linked Antenatal and Newborn Screening Programme*, available on the National Screening programme website.

**Interfaces:**

- The referral of pregnant women presenting through primary care into midwifery services, or directly to a prenatal diagnosis centre if appropriate
- The sending of blood sample and FOQ (ideally electronically) from the midwifery service to the antenatal screening laboratory
- The sending of results from the antenatal screening laboratory back to the midwife (ideally using antenatal status codes)
- The referral of women/couples with confirmed carrier or affected status for counselling by the midwifery service
- The referral of women/couples with confirmed carrier or affected result to prenatal diagnosis centre by the midwifery service
- The reporting of results from the prenatal diagnosis centre to the referring clinician
- The informing of parents, community midwife and GP of prenatal diagnosis testing results by the referring clinician
- Maternity unit and newborn laboratory (maternity services and newborn sickle cell screening also covered by NHS Newborn Blood Spot Screening Programme). This may include
  - Midwife notifies a new birth and NHS number is issued, automatic notification to local CHRD
  - Midwife responsible for care sends blood spot card to newborn screening laboratory
  - Failsafe system to ensure laboratory receipt of sample
  - Laboratory requests midwifery services for a repeat if required
  - Laboratory sends results to CHRD, ideally electronically
  - CHRD checks for untested babies within effective timeframe
  - Laboratory refers screen positive results to specialist teams
  - CHRD send normal results letter to health visitor and to parents
  - CHRD informs maternity or health visiting services of untested babies
  - CHRD use local pathway for reporting carrier results
Public health functions to be exercised by NHS England

- Clinician informs CHRD if unable to complete screen so it can be recorded on the baby’s record
- Health visitors ensure parents receive results and record results in PCHR by 8 weeks

- Newborn laboratory and care services. This should be guided by The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services

The Map of Medicine describes these interfaces in more detail and can be found at: [http://eng.mapofmedicine.com/evidence/map/index.html](http://eng.mapofmedicine.com/evidence/map/index.html)

3.9 Information on test/screening programme

In accordance with UK NSC standards and protocols the provider will ensure that during pregnancy and after birth parents/carers are provided with approved information on sickle cell and thalassaemia screening by a trained workforce. The provider will ensure women have seen the written information *Screening tests for you and your baby* or had access to it in an appropriate format.

In addition, a wide range of information for the SCT Programme has been developed in a range of formats and media for women/couples that is available for local use.

3.10 Testing (laboratory service, performance of test by individuals)

Antenatal, newborn and DNA referral laboratories are expected to follow the policy guidance and standards laid out in ‘Sickle Cell and Thalassaemia: Handbook for Laboratories’ and meet the programme standards.

3.11 Results reporting and recording

Antenatal results

- Antenatal laboratory sends results as per local arrangements, ideally using antenatal status codes
- The maternity unit notifies the newborn screening laboratory of carrier status
- It is recommended that the maternity unit keeps a log of and notifies the newborn screening laboratory of all women who are carriers and affected as well as carrier couples
Newborn results

- Newborn laboratory sends results to CHRD, ideally electronically using nationally approved status codes
- CHRD record conclusive results on a child health information system for all the eligible population
- CHRD informs maternity or health visiting services of null/incomplete results
- Clinician informs CHRD if unable to complete screen so it is documented in the baby’s record
- As systems are developed it is anticipated that newborn screening results should routinely be transferred to primary care in a standard format.
  - Screen positive results are reported according to screening programme standards and care commissioning guidelines (The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services)

3.12 Results giving

In accordance with UK NSC standards and protocols the provider will ensure that a healthcare professional informs parents of their results.

For newborn screening results:

- the CHRD will send a normal results letter to parents and notify the health visitor
- the health visitor will ensure that parents receive the results and record the results in the Personal Child Health Record by 8 weeks
- sickle cell carrier (and specified haemoglobin variant) results should ideally be given face to face by a trained healthcare professional
- all screen positive results will be given to parents by a trained health professional face-to-face by four weeks of age, following local protocols and ensure that the baby enters care by eight weeks of age

3.13 Transfer of and discharge from care obligations

Based on the generic screening objectives of the programme the antenatal screening pathway ends for:

- pregnancies resulting in a live birth when the antenatal screening results are included on the blood spot card (This includes screening and PND results) (Node 27 on map of medicine)
- women opting for termination of pregnancy when the woman is counselled appropriately following prenatal diagnosis (nodes 26,27 and 28 on map of medicine).
Based on the generic screening objectives of the programme the newborn screening pathway ends for:

- Condition not suspected and carrier results when parents and GPs are informed of the result (node 22 /23 on map of medicine)
- Screen positive results when the parents are informed of the result and the baby is seen and tested and diagnosis confirmed by a clinician and registered in the designated clinical network (node 24 on map of Medicine). More detail is available in The National Haemoglobinopathies Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services

3.14 Patient/carer information

A variety of national patient information materials are available from the SCT National Screening programme website.

Educating patients and carers to come to terms with their disease and how to best manage their condition is included within

3.15 Exclusion criteria

See Section 3.4 for details of exclusion criteria in the antenatal period.

Newborn:
- Babies stillborn or who died before day 8
- Children over 1 year of age

3.16 Staffing

In accordance with UK NSC standards and protocols the provider will ensure that there are adequate numbers of appropriately trained staff in place across the screening pathway to deliver the screening programme in line with best practice programme and laboratory guidelines. Qualifications will be specific to staff delivering the service across the care pathway. Staff must demonstrate competence (which is linked to training).

The Provider will have in place a workforce plan designed to maintain a sustainable programme, especially where increase in birth rate are predicted and/or where there are difficulties in the recruitment of appropriately qualified healthcare staff.
All professionals involved in the screening programme are required to keep up to date with nationally approved training programmes.

The provider will ensure that counsellors for the sickle cell and thalassaemia screening programme are trained in an approved course see (sct.screening.nhs.uk/externaltraining).

3.17 User involvement

In accordance with UK NSC standards and protocols the provider will be required to:

- demonstrate that they have collected (or have plans in place to collect) the views of service users, families and others in respect of the services they provide
- demonstrate how those views will influence service planning and delivery for the purposes of raising standards
- show that all families are given information about how to provide feedback about services they receive, including about the complaints procedure.

Collection of the views of service users/families will often be via surveys or questionnaires. It is expected that such surveys will take place on a regular (rather than ad hoc) basis and that the results will be made available to NHS England on request.

3.18 Premises and equipment

In accordance with UK NSC standards and protocols the provider will ensure that:

- suitable premises and equipment are provided for the screening programme
- appropriate policies are in place for equipment calibration, maintenance and replacement
- appropriate IT systems are in place to support programme delivery including audit and monitoring functions and developing electronic links with other Providers in the screening pathway
- the UK NSC Laboratory Guidelines Sickle Cell and Thalassaemia: Handbook for Laboratories’ are followed. These can be found on the SCT National Screening programme website
Safety & Safeguarding

The provider should refer to and comply with the safety and safeguarding requirements as set out in the NHS Standard Contract. As an example, please see link below for 2013/14 NHS Standard Contract:

Section 4: Service Standards, Risks and Quality Assurance

4.1 Key criteria and standards

Programme standards are available on the programme website (http://sct.screening.nhs.uk/standards). Providers will meet the acceptable and work towards the achievable programme standards. A number of resources to support providers are available on the programme website. See antenatal, newborn and linked SCT standards found in “NHS Sickle Cell and Thalassaemia Screening Programme: Standards for the linked Antenatal and Newborn Screening Programme”.

The SCT National Screening programme supports health professionals to meet these standards and deliver a high quality SCT screening programme. A number of resources to support health professionals are available on the SCT National Screening programme website.

4.2 Risk assessment of the screening pathway

Providers are expected to have an internal quality assurance and risk management process that assures the commissioners of its ability to manage the risks of running a screening programme.

Providers will:

• ensure that appropriate failsafe mechanisms are included across the whole screening pathway
• review and risk assess local screening pathways in the light of guidance offered by Quality Assurance processes or the National Screening programme
• work with the Commissioner and Quality Assurance Teams to develop, implement, and maintain appropriate risk reduction measures
• ensure that mechanisms are in place to regularly audit implementation of risk reduction measures and report incidents
• ensure that appropriate links are made with internal governance arrangements, such as risk registers
• ensure routine staff training and development is undertaken
On a quarterly basis high scoring risks will be identified and agreed between the provider and the commissioners and plans put in place to mitigate against them. Risk identification should take into account failsafe mapping (please also see section 2.3 Failsafe).

4.3 Quality assurance

Providers will participate fully in national Quality Assurance processes and respond in a timely manner to recommendations made. This will include the submission to QA teams and commissioners of:

- data and reports from external quality assurance schemes
- minimum data sets as required – these may be required to be submitted to national external bodies e.g. National Vascular Database etc.
- self-assessment questionnaires / tools and associated evidence
- audits or data relating to nationally agreed internal quality assurance processes

Providers will participate fully in the QA visit process where required and cooperate in undertaking ad-hoc audits and reviews as requested.

Providers will respond to QA recommendations by the submission of action plans to address identified areas for improvement and any non-conformities / deviations from recommended performance thresholds.

Where QA believe there is a significant risk of harm to the population, they will recommend to commissioners to suspend a service.

Laboratories undertaking sickle cell and thalassaemia screening should

- be accredited by CPA or equivalent and list the screening tests in their repertoire of services ([http://www.cpa-uk.co.uk/](http://www.cpa-uk.co.uk/))
- participate in and respond in a timely manner to an accredited external quality assurance scheme for sickle cell and thalassaemia screening. e.g. UKNEQAS scheme
- Make available timely data and reports from external quality assurance programmes and accreditation services to screening programmes, national teams and commissioner
4.4 Serious incidents

Providers will comply with the national guidance for the management of incidents in screening programmes and NHS England guidance for the management of incidents.

“Managing Incidents in England NHS National Screening Programmes Interim Guidance”


4.5 Procedures and Protocols

The provider will be able to demonstrate that they have audited procedures, policies and protocols in place to ensure best practice is consistently applied for all elements of the screening programme.

4.6 Continual service improvement

Where national recommendations and acceptable/achievable standards are not currently fully implemented the provider will be expected to indicate in service plans what changes and improvements will be made over the course of the contract period.

The provider shall develop a CSIP (continual service improvement plan) in line with the KPIs and the results of internal and external quality assurance checks. The CSIP will respond and any performance issues highlighted by the commissioners, having regard to any concerns raised via any service user feedback. The CSIP will contain action plans with defined timescales and responsibilities, and will be agreed with the commissioners.

4.7 Teaching and training

The provider will ensure that:

- Education, training and staff development are an integral part of the service and complies with the requirements of the screening programme
- It keeps up to date with clinical advances
- Contributes to education and training of other relevant professionals where appropriate
It should also aspire to participate in properly conducted quality research where possible (with appropriate ethical approval).
Section 5: Data and Monitoring

5.1 Data collection, monitoring and reporting

Activity, performance and KPI data will be collected by providers and shared with NHS ENGLAND to allow benchmarking between areas within the eligible screening programme population.

5.2 Key performance indicators / Public Health Outcomes Framework

The provider shall adhere to the requirements specified in the document ‘Key Performance Indicators for Screening. Please refer to http://www.screening.nhs.uk/kpi for further details, guidance and updates on these indicators.

Public Health Outcomes Framework Indicator

2.21iii: The percentage of pregnant women eligible for antenatal sickle cell and thalassaemia screening for whom a conclusive screening result is available at the day of report

Key Deliverable: The acceptable level should be achieved as a minimum by all services

Acceptable ≥ 95.0%
Achievable ≥ 99.0%
2012-13 national baseline is 98%

Providers to implement an additional KPI on coverage of untested babies (movers-in) up to one year of age.

This section outlines what data (in addition to Key Performance Indicators) the provider will be expected to collect and submit. Please refer to http://sct.screening.nhs.uk/datacollection for further details, guidance and updates. There are two main streams of the data collection process:
Routine reporting:

- Standard annual data returns by:
  (i) all antenatal laboratories,
  (ii) all newborn laboratories
- (iii) the three DNA laboratories (prenatal diagnosis) including data on pregnancy outcomes for those women who have undergone prenatal diagnosis.
- (iv) anonymous data from designated clinical centres (see The National Haemoglobinopathy Project: A Guide to Effectively Commissioning High Quality Sickle Cell and Thalassaemia Services). These data include for example:- timely referral to care, confirmed screening result and information about vaccination status.
- (v) two DNA laboratories (newborn screening) for babies who have had a blood transfusion
- SCT antenatal KPIs are submitted from maternity units and antenatal laboratories to the UK NSC quarterly and annually (submission of KPI data began in Q4 of 2010/11).

Other:

- Programme evaluation data collection with support from the Health Research Authority Confidentiality Advisory Group.
- Ad hoc surveys to inform screening pathway and processes
- Incident reporting

There is on-going work under development about data collection.