Other Disorders
Hemoglobinopathies

A hemoglobinopathy is a condition that affects the red blood cells and results from genetically determined changes in the molecular structure of hemoglobin. In the laboratory, the hemoglobin Isoelectric Focusing (IEF) and High-Performance Liquid Chromatography (HPLC) tests will reveal multiple hemoglobinopathy disorders with varying degrees of severity.

Their effects range from mild anemia in Hemoglobin C disease (Hemoglobin CC) and C, Beta (β) Thalassemia, to severe pain episodes, growth delays, increased susceptibility to infections and persistent anemia in Sickle Cell Anemia (Hemoglobin SS) and S, β Thalassemia.

Hemoglobinopathies are inherited in an autosomal recessive pattern. Carriers of a single abnormal gene for these disorders are considered to have a trait. Persons with a trait will have red blood cells that contain a mixture of normal and abnormal hemoglobin. Most hemoglobinopathy traits cause no disease or anemia under normal physiologic conditions*.

| Inheritance: | Autosomal recessive |
| Estimated Incidence: | 1:400 African Americans (sickling disorders)  
1:2500 All Races/Ethnicities (sickling disorders) |
| Neonatal Presentation: | None |
| Method of Notification: | All abnormal results are called and faxed to the provider of record. |
| Next Steps if Abnormal: | **Sickling disorders** - Refer to pediatric hematologist. Consider initiation of penicillin prophylaxis upon receipt of newborn screening report if the hemoglobin pattern is FS, FSA, FSB, FSC, FSE/O or FSD/G or FSV, **Report all subsequent findings to state newborn screening program.**  
**Hemoglobin C Disease or β Thalassemia** - Refer to a pediatric hematologist. **Report all findings to state newborn screening program.**  
If all other newborn screening results are normal, a repeat specimen newborn screen is not required. Initial sample will be sent by the lab for hemoglobin confirmation.  
All hemoglobinopathies and traits - Refer family to a sickle cell foundation for family testing and counseling. |
| Abnormal Screen Result: | See Chart below |
The following table outlines retesting procedures for the most common results of the screen. It is important to remember that PREMATURITY AND TRANSFUSIONS AFFECT TEST RESULTS. Each type of hemoglobin in the infant's blood is identified by a letter on the test result (e.g. F=Fetal, A=Adult or normal, S=Sickle, V=unknown variant).

The position of the letter represents the amount of hemoglobin type present with the hemoglobin of greatest concentration listed first. (Example: "FSA" usually indicates a sickling disorder and "FAS" indicates a trait).

When rare a hemoglobin is detected, specific instructions will be sent from CH. A portion of the abnormal bloodspot will also be sent to the Children’s Hospital of Oakland Research Institute (CHORI) for confirmatory testing. If all other newborn screening results are normal, a repeat specimen is not required.

<table>
<thead>
<tr>
<th>Newborn’s Hemoglobin Result</th>
<th>Potentially indicative of:</th>
<th>Sent to CHORI?</th>
</tr>
</thead>
<tbody>
<tr>
<td>FA</td>
<td>Normal Newborn Hemoglobin</td>
<td>No</td>
</tr>
<tr>
<td>AF</td>
<td>Normal or transfused hemoglobin</td>
<td>No</td>
</tr>
<tr>
<td>FS</td>
<td>Sickle Cell disease, Sickle β0-thalassemia or Sickle with Hereditary Persistence of Fetal Hemoglobin (S-HPFH)</td>
<td>Yes</td>
</tr>
<tr>
<td>FSA</td>
<td>Sickle β+-thalassemia or Sickle cell trait</td>
<td>Yes</td>
</tr>
<tr>
<td>FSB (FS + Bart’s)</td>
<td>α Thalassemia with Sickle Hemoglobin</td>
<td>Yes</td>
</tr>
<tr>
<td>FSC</td>
<td>Sickle C disease, SC Harlem</td>
<td>Yes</td>
</tr>
<tr>
<td>FSD</td>
<td>Sickle D Disease</td>
<td>Yes</td>
</tr>
<tr>
<td>FSE</td>
<td>Hemoglobin SE Disease</td>
<td>Yes</td>
</tr>
<tr>
<td>FSG</td>
<td>Sickle Cell Anemia, Sickle cell β Thalassemia, Sickle G Philadelphia</td>
<td>Yes</td>
</tr>
<tr>
<td>FSO</td>
<td>Sickle O Arab Disease</td>
<td>Yes</td>
</tr>
<tr>
<td>FSV</td>
<td>Sickle with Variant Hemoglobin pattern</td>
<td>Yes</td>
</tr>
<tr>
<td>FC</td>
<td>Homozygous Hemoglobin C disease or Hemoglobin C β0 thalassemia</td>
<td>Yes</td>
</tr>
<tr>
<td>FCA</td>
<td>Hemoglobin C β+ thalassemia or Hemoglobin C trait</td>
<td>Yes</td>
</tr>
<tr>
<td>FCE</td>
<td>Hemoglobin CE Disease</td>
<td>Yes</td>
</tr>
<tr>
<td>FCV</td>
<td>Hemoglobin C Variant</td>
<td>Yes</td>
</tr>
<tr>
<td>FDD</td>
<td>Homozygous Hemoglobin D, Hemoglobin D Thalassemia</td>
<td>No</td>
</tr>
<tr>
<td>FDA</td>
<td>Hemoglobin D/β Thalassemia or Hemoglobin D trait</td>
<td>No</td>
</tr>
<tr>
<td>FDV</td>
<td>Hemoglobin D Disease, Hemoglobin D Thalassemia, or Hemoglobin D trait</td>
<td>No</td>
</tr>
<tr>
<td>FE</td>
<td>Homozygous Hemoglobin E Disease, Hemoglobin E β+ thalassemia or Hemoglobin E β0 thalassemia</td>
<td>Yes</td>
</tr>
<tr>
<td>Code</td>
<td>Description</td>
<td>Value</td>
</tr>
<tr>
<td>------</td>
<td>-----------------------------------------------------------------------------</td>
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<tr>
<td>FEA</td>
<td>Hemoglobin E β+ thalassemia or Hemoglobin E trait</td>
<td>Yes</td>
</tr>
<tr>
<td>FEV</td>
<td>Hemoglobin E Disease, Hemoglobin E β+ thalassemia, Hemoglobin E β0 thalassemia or Hemoglobin E trait</td>
<td>Yes</td>
</tr>
<tr>
<td>FV</td>
<td>Unknown hemoglobin variant</td>
<td>No</td>
</tr>
<tr>
<td>FO</td>
<td>Homozygous Hemoglobin O-Arab</td>
<td>No</td>
</tr>
<tr>
<td>FVA</td>
<td>Unknown hemoglobin variant</td>
<td>No</td>
</tr>
<tr>
<td>FOA</td>
<td>Hemoglobin O-Arab/β+ Thalassemia or Hemoglobin O-Arab/β0 Thalassemia</td>
<td></td>
</tr>
<tr>
<td>FF</td>
<td>Premature Infant, Hereditary Persistence of Fetal Hemoglobin (HPFH) or Homozygous β thalassemia major</td>
<td>Yes</td>
</tr>
<tr>
<td>FAB*</td>
<td>Hemoglobin Bart’s - α thalassemia of unknown severity to Hemoglobin H disease</td>
<td>No</td>
</tr>
<tr>
<td>FAC, FAD, FAE, FAG, FAO, FAS, FAV, FA + fast band, FAB &lt; 15% (Bart’s)</td>
<td>Various Hemoglobin traits/carriers</td>
<td>No</td>
</tr>
</tbody>
</table>

Please consult with a pediatric hematologist for further recommendations.

Treatment: **Sickling disorders** - Penicillin/antibiotic prophylaxis beginning in infancy and continuing through early childhood. Prompt evaluation/management of acute illness to lessen development of sickling crisis, particularly if fever is present.

Appropriate pain management strategies (such as use of extra fluids, oral analgesics, comfort measures), including rapid triage if home management strategies are not sufficient.

Transfusion may be necessary in certain instances. Medications to increase the production of fetal hemoglobin and lower leukocyte counts such as hydroxyurea may be used in certain children.

A blood or marrow transplant is the only known cure for sickle cell disease (SCD). However, transplant has serious risks and is only used in patients with severe SCD who have symptoms including stroke, acute chest syndrome, and frequent pain episodes. The transplant replaces diseased blood-forming cells with healthy ones.

The type of transplant used to treat SCD is an allogeneic transplant. This type of transplant uses healthy blood-forming cells from a family member, unrelated donor, or umbilical cord blood unit.

For an allogeneic transplant, a patient gets chemotherapy, with or without radiation, prior to transplant to prepare his or her body for the treatment. Then, the replacement cells are infused
into the patient’s blood stream. From there, the cells find their way into the bone marrow, where they start making healthy white blood cells, red blood cells and platelets.

The entire process, from the start of chemotherapy or radiation until hospital discharge, can last weeks to months followed by many months of recovery at home.

**Special Considerations**

**Transfusion** - Transfusion of red blood cells prior to drawing the newborn screening specimen will affect the hemoglobinopathy result. Repeat screening for hemoglobinopathies should be done 120 days after the last transfusion. If the date of the last transfusion is unknown, put the date of hospital discharge on the collection form next to “Transfused”.

**Specimen Analysis at the Reference Laboratory** - The initial newborn screening bloodspots for infants with hemoglobinopathy results indicative of disease are sent to the Children’s Hospital of Oakland Research Institute (CHORI) for more specific hemoglobinopathy analysis and genetic testing. The result of the CHORI analysis is sent to the provider of record upon receipt by the Laboratory.

**Special Follow-up Assistance** - The DHEC Children and Youth with Special Healthcare Needs (CYSHCN) Sickle Cell Program assists primary care providers across the four regions of the state to ensure infants identified with a sickling disorder are seen by a pediatric hematologist within the first six weeks of age. They coordinate activities with pediatric hematologists, Sickle Cell Foundations, health departments and hospitals, so that families are directed to the services closest to them.

**Participation in Sports or Extreme Physical Activity** - Some persons with sickle cell trait may exhibit a sickling crisis associated with extreme physical activity. Precautions must be taken to lessen the chance for exertional rhabdomyolysis.

**Coordination of Care** - In coordination with SC DHEC Children and Youth with Special Healthcare Needs (CYSHCN) Sickle Cell Program and the Sickle Cell Foundations of South Carolina, counseling, education and other resources are offered to families of children diagnosed with a hemoglobin disorder and those with children carrying a mutation identified through newborn screening.

The goals of counseling are to increase the understanding of genetic diseases, discuss disease management options, and explain the risks and benefits of testing. Counseling sessions focus on giving vital, unbiased information and non-directive assistance in the patient's decision-making processes.
Sickle Cell Foundation Contacts in South Carolina

Community Based Organizations (CBO’s) for Support:

COBRA Human Services Agency Sickle Cell Program
3962 Rivers Ave
PO Box 71473
Charleston, SC 29415
*Toll Free* (800) 354-4704
(843) 225-4866, Service Line
(843) 225-4869, Fax
cobraagency@bellsouth.net

Orangeburg Area Sickle Cell Foundation
825 Summers Ave
PO Box 892
Orangeburg, SC 29116
(803) 534-1716, Phone
(803) 531-2422, Fax
orangeburgsickle@aol.com

James R. Clark Memorial Sickle Cell Foundation
1420 Gregg St
Columbia, SC 29201
*Toll Free* (800) 506-1273
(803) 765-9916, Phone
(803) 799-6471, Fax
www.jamesrclarksicklecell.org
office@jamesrclarksicklecell.org

Louvenia Barksdale Sickle Cell Anemia Foundation
645 S Church St
PO Box 191
Spartanburg, SC 29304
(864) 582-9420, Phone
(864) 582-9421, Fax
www.barksdalesicklecell.org
ldbarksdale@charter.net

Centers for Disease Control and Prevention
Sickle Cell Disease (SCD) National Resource Directory
https://www.cdc.gov/ncbddd/sicklecell/index.html
Cystic Fibrosis (CF)

Cystic fibrosis (CF) is characterized by pulmonary obstruction often accompanied by exocrine pancreatic dysfunction. A defect in the cystic fibrosis transmembrane conductance regulator (CFTR) gene leads to obstruction of exocrine pancreatic ducts, which causes an increase in the pancreatic enzyme immunoreactive trypsinogen (IRT) in blood.

CF usually affects the lungs, pancreas, intestines, liver and sweat glands, causing failure to thrive, steatorrhea, intestinal obstruction, salt loss, and progressive obstructive lung disease.

Inheritance: Autosomal recessive

Estimated Incidence: 1:3,900 (varies by ethnic group)

Abnormal Screen Result: Elevated IRT

Method of Notification: All abnormal results are called and faxed to the provider of record and reflexed to CF 2nd tier testing. Abnormal lab reports are also faxed to the regional pulmonologist where the baby is located.

Next Steps if Abnormal: If initial IRT is elevated and no mutations are found on CF 2nd tier test, see infant to ascertain health status. Repeat IRT on filter paper as soon as possible.

If repeat IRT is within normal limits, no further bloodspots are needed.

If IRT is still elevated on repeat testing or mutations are found on CF 2nd tier test, consult pediatric pulmonologist for further instructions and evaluation.

Diagnosis by sweat chloride testing at a CF Foundation accredited care center is necessary for final diagnosis. Initiate treatment as recommended by specialist.

Report all findings to state newborn screening program.

Neonatal Presentation: Usually none. Meconium ileus or volvulus may occur in 5-10 % of affected infants. Prolonged jaundice without other cause is more common than very early lung disease.

Treatment: Chest physiotherapy to aid in airway clearance. Antibiotics/other medications to treat lung infections as needed. Pancreatic enzymes if indicated; vitamins; NaCl supplements. Close monitoring of growth parameters and use of nutritional supplements if needed to enhance/maintain appropriate growth/development.
Special Considerations

Premature/Sick Infants - The stress of prematurity and/or illness can lead to falsely elevated IRT test results.

Meconium Ileus - All infants with meconium ileus should be thoroughly evaluated for CF regardless of the IRT result. A normal IRT result does not rule out CF in these infants.

Prenatal Screening and confirmatory testing - For general population CF carrier screening, the American College of Medical Genetics (ACMG) and American College of Obstetricians and Gynecologists (ACOG) recommend a core panel of 23 mutations that will identify 49–98% of carriers, depending on ethnic background.

The SC DHEC Public Health Laboratory will perform an extended confirmatory panel of 60+ mutations for screen positive infants. The extended panel includes the recommended core panel of 23 mutations, thereby ensuring comprehensive mutation coverage.

However, negative carrier status in the infant or parents does not definitively rule out the possibility of CF in an infant. Infants may have other rare mutations that are not included in a standard CF 2nd tier test.

False Negative Test Results - Some infants with CF may have false negative IRT results.

Physicians must remain alert to clinical signs of CF in older infants despite normal initial screening results.
Severe Combined Immunodeficiency (SCID)

Low levels of T-cell Receptor Excision Circles (TRECs) are associated with Severe Combined Immunodeficiency (SCID). Other conditions associated with low TRECs include reticular dysgenesis, coronin-1A deficiency and thymic aplasia/complete DiGeorge syndrome. T lymphocytes fail to develop and the affected infant may also have impaired B lymphocyte function.

Inheritance: Autosomal recessive and X-linked

Estimated Incidence: 1:40,000 to 1:60,000

Abnormal Screen Result: Elevated Cq

Method of Notification: All abnormal results are called to provider of record and the Immune Disorder Specialist

Next Steps if Abnormal: Potential medical emergency when TRECs are low and RNase P is within normal limits! The screening report will indicate Cq (Quantification Cycle) value instead of actual number of TRECs. Cq is the number of test cycles needed for the fluorescence of the amplified DNA to exceed the laboratory’s established fluorescence threshold.

The Cq value of TRECs is inversely related to the copy number of TRECs in a specimen. Specimens that have a low TREC content (low copy number) have a higher Cq value.

See infant as soon as possible to ascertain health status. Consult pediatric specialist (immunology or pediatric infectious disease) and initiate diagnostic evaluation and treatment as recommended. Common diagnostic studies include specialized flow cytometry and molecular testing to determine specific mutations. Report all findings to state newborn screening program.

In addition, repeat TREC on filter paper and send to the DHEC laboratory. Low TRECs with low RNase P may indicate DNA amplification failure. Prompt repeat screening is necessary to rule out SCID in these infants.

Neonatal Presentation: Usually none. Median age for onset of symptoms is 8 weeks of age.

Emergency Treatment: Usually none.

Standard Treatment: Bone marrow transplantation by 3 months of age is associated with the best outcomes for SCID. Infants with other conditions may be treated with medications.
**Special Considerations**

**Infectious Disease Precautions** - Parents should be instructed to avoid exposure of the infant to anyone with viral/bacterial illnesses until SCID is definitively ruled out. No vaccines should be given until cleared to do so by the specialist.

The specialist may advise breastfeeding mothers to suspend breastfeeding while their blood is checked for anti-CMV IgG antibodies and CMV DNA. These mothers should be encouraged to pump and freeze their breast milk during this time. Prompt resumption of breastfeeding is encouraged if the mother is seronegative.

**Only leukoreduced, CMV negative, irradiated blood should be used if a transfusion is necessary.**

**Premature/Sick Infants**—Premature infants may have low TREC due to immaturity of the immune system. Prompt repeat screening is indicated. The pediatric specialist (immunology or pediatric infectious disease) may recommend flow cytometry if TREC are low in a second blood spot specimen.

Low TREC may also be found in specimens obtained from infants who have undergone thymectomy/cardiac surgery if the specimen is collected after surgery.
Hearing Loss (HL) and Critical Congenital Heart Defects (CCHD)*

*These point of care newborn screening tests (not blood tests) are administered at the hospital or other birthing facility.

For newborn hearing screening and hearing loss information, please refer to SC DHEC First Sound Hearing Screening Program. For CCHD information refer to the SC DHEC Birth Defects Program.

First Sound Program Manager/Audiologist:
Tara Carroll, MCD, CCC/A.................................803-898-0708
email: carroltp@dhec.sc.gov

Birth Defects Program Manager:
Vinita Oberoi Leedom, MPH, CIC..........................803-898-0771
email: leedomvo@dhec.sc.gov