Parents’ Guide to Sickle Cell Hemoglobin E Disease (SE Disease)

You have just learned that your baby has sickle cell hemoglobin E disease (Hemoglobin Hb SE). The information below will help answer some of your questions. However, it should not take the place of an informed discussion with your baby’s doctor (primary care provider).

What is Hemoglobin?
Hemoglobin is the main ingredient in red blood cells. Hemoglobin helps red blood cells carry oxygen from the lungs to all parts of the body and gives blood its red color. There are many hemoglobin types (this is not the same as a blood type). The main hemoglobin type in normal red blood cells is hemoglobin A. The presence of hemoglobin A in red blood cells makes them smooth and shaped with a large, rounded depression on each side of the cell, known as a biconcave disk shape (see picture left above). This shape allows normal red blood cells to be very flexible, moving easily through the blood vessels to deliver oxygen to the body.

What is Sickle Cell Hemoglobin E Disease (Hemoglobin SE)?
Hemoglobin SE disease (Hemoglobin SE) is a type of sickle cell disease. People with Hemoglobin SE disease have red blood cells that contain both hemoglobin S (sickle hemoglobin) and hemoglobin E. These individuals do not have hemoglobin A. People inherit this disease from both parents. This is called an autosomal recessive genetic disorder. This means a person with Hemoglobin SE disease inherited hemoglobin E from one parent and hemoglobin S from the other parent. There is a 1 in 4 or 25% chance with each pregnancy of having a child with Hemoglobin SE disease. These are the possible outcomes with each pregnancy (see picture).

- 25% (1 in 4) chance of having a child with hemoglobin E trait* (A/E)
- 25% (1 in 4) chance of having a child with sickle cell trait* (A/S)
- 25% (1 in 4) chance of having a child with hemoglobin SE disease (S/E)
- 25% (1 in 4) chance of having a child with hemoglobin A (normal) (A/A)

*People with hemoglobin E trait or sickle cell trait do not have a hemoglobin disease. They cannot develop disease later in life. They can pass the trait on to their children.

Hemoglobin SE disease is found in males and females equally and occurs in all races. Sickle hemoglobin is most common among people of African, Caribbean and South American ancestry. It is less common in Mediterranean and Middle Eastern people. Hemoglobin E occurs most often among people of Southeast Asian origin. All newborn babies in Utah are tested for hemoglobin SE disease, regardless of their race or ethnic background. Hemoglobin SE disease is a lifelong condition and is not contagious.

What Problems can Sickle Cell E Disease Cause?
Hemoglobin E interacts with hemoglobin S. In certain situations hemoglobins may sickle. Sickling is a condition in which the red blood cells become hard and sticky and clog up small blood vessels to the lungs, bones or other body tissues. This is similar to situations when hemoglobin S interacts with another S, C, D or O hemoglobin. Those with SE disease may have periods of feeling well and times of sudden intense pain that can be anywhere in the body. Damaged red blood cells cannot deliver oxygen to the body and over time may damage the body’s tissues and organs.

Hemoglobin SE red blood cells are not very flexible in moving through the blood vessels. The sickle E (SE) red blood cells sickle and rupture. They have a life span that is shorter than normal red blood cells. (Normal blood cells live about 120 days). This leads to anemia (low red blood cell count) and decreases the ability of red blood cell to hold onto oxygen.
Bilirubin is produced from the remains of the ruptured cells which, in turn, can cause jaundice (yellowing of the skin). Children with sickle cell (SS) disease in general sometimes may have serious health problems that, if not treated, can lead to death. Children with sickle cell hemoglobin E (SE) disease, however, are expected to have a milder course than those with SS or SC disease. These problems are: infection of the blood (septicemia) and sudden enlargement of the spleen with a drop in the red blood cell count due to trapping of a large amount of blood in the spleen (acute splenic sequestration). Other serious problems include: sudden pain, swelling of hands and feet, fever, increased infections, anemia, chest pain and trouble breathing, pneumonia, stroke, blood in urine, leg ulcers, gallstones, vision problems, organ damage, kidney failure, painful erections, and problems during pregnancy.

What can be done to Treat Sickle Cell E Disease?
If your child who has been diagnosed with sickle cell hemoglobin E disease (SE) is having any of the symptoms listed above, consult your child's doctor for instructions.

- **Pain Management.** Recommended home treatment includes applying a heating pad to the painful area and medicines such as acetaminophen (Tylenol®) or ibuprofen (Advil® and Motrin®). Drinking plenty of liquids and rest can help. Once the pain goes away children are usually active again. If the pain is not better after taking medicine at home, your child may need treatment with stronger medicine in the emergency room or to be admitted to the hospital for treatment.

- **Penicillin.** Very serious infections of the blood sometimes occurs in infants and young children with hemoglobin SE cell disease. Penicillin prescribed by your child's doctor taken two times each day can help decrease these infections in your child. Keeping your child’s immunizations up-to-date will significantly decrease the chance of getting severe infections.

What are the Most Important Things to Remember?
- Work closely with your child's doctor and hematologist (a doctor who specializes in blood disorders). Make sure your child has regular checkups and call your child’s doctor with questions.
- Your child should avoid extreme hot and cold temperatures. He/she should avoid becoming exhausted, get plenty of rest, and drink lots of water to reduce chances of having pain.
- Check your child’s spleen as advised by the hematologist. Your child’s doctor will show you where the spleen is and what feels normal. If the spleen suddenly feels larger, your child should be seen as soon as possible by his or her doctor.
- Your child will need an immediate medical evaluation for a fever of 38.5°C (101°F) or greater, difficulty breathing or chest pain. Take your child to a facility that provides emergency care.
- If your child has no energy and looks very pale he or she should be evaluated by a doctor.
- Be sure your child receives all childhood immunizations when they are scheduled and any additional immunizations recommended by your child’s doctor.
- Call your child's doctor if you have questions and have your child seen if you have medical concerns.

What Additional Treatment are Available?
Blood transfusions and medications to decrease or prevent the formation of sickle-shaped red blood cells may be used. Those with severe hemoglobin SE disease may be treated by a bone marrow transplant from a compatible donor, but this is considered a high-risk procedure.

How Do I Get More Information?
Talk with your baby’s doctor. We recommend that you make an appointment with a pediatric hematologist in the near future. You may also want to have a genetic consultation for you and your family to see how hemoglobin SE disease might affect future children or grandchildren.