Sickle cell disease, part 1: Understanding the condition

Sickle cell disease (SCD) is one of the most common genetic blood disorders in the world and affects approximately 12,500 individuals in the UK. This article looks at the definition of SCD, symptoms and treatments, and practical tips to prevent sickle cell crises.

Sickle cell disease (SCD) is a group of genetic disorders that affect haemoglobin. This is the molecule in red blood cells that delivers oxygen to cells throughout the body. People with SCD have atypical haemoglobin molecules called haemoglobin S, which can alter red blood cells into a sickle or crescent shape (Serjeant and Serjeant, 2001). In Britain it is most common in people of African or Afro-Caribbean descent but may also occur in people from India, Pakistan, the Middle East or eastern Mediterranean. An estimated 12,500 individuals live with SCD in England (Sickle Cell Society, 2008).

Genetic inheritance
Sickle cell disease is an autosomal recessive genetic disorder. For the disease to manifest a person must inherit two copies of the sickle cell gene—one from each parent. People who inherit a sickle cell gene from one parent and a normal gene from the other have a condition called sickle cell trait (Hb AS). They do not have the disease but they possess one of the genes that causes it and can therefore pass this gene to their children. For example, if one parent has sickle cell anaemia and the other is normal all of the children will have sickle cell trait. When both parents have sickle cell trait they have a 25% chance of having a baby with sickle cell disease at every pregnancy (Serjeant and Serjeant, 2001; Oni et al, 2006).

How does sickling happen?
Normal red cells are biconcave, soft and flexible. They maintain their shape as they pass through the capillaries and release oxygen to the peripheral tissues. However, when sickle haemoglobin releases oxygen in the peripheral tissues the molecules tend to stick together and form long chains or polymers. The polymer is a rope-like fibre that aligns with others to form a bundle distorting the red cell into classic crescent or sickled forms. The problem is not simply one of abnormal shape as the deformed cells are rigid. They block the flow of cells and interrupt the delivery of oxygen to the tissues causing ischemia, pain and tissue damage (Bain, 2002; Stuart and Nagel, 2004) (Figure 1).

Types of sickle cell disease
There are different types of sickle cell disease. The most common are Haemoglobin SS or sickle cell anaemia, Haemoglobin SC disease, and Haemoglobin sickle beta-thalassaemia (Table 1).

Thalassaemia genes produce normal haemoglobin but in variably reduced amounts. If the gene produces no normal haemoglobin, which is sickle beta zero thalassaemia, the condition is virtually identical to sickle cell anaemia. Some patients have a gene that produces a small amount of normal haemoglobin, which is sickle beta plus thalassaemia, that tends to result in a mild form of sickle cell disease. Sickle beta-thalassaemia is the most common sickle syndrome seen in people of Mediterranean descent such as Italians, Greeks and Turks (Franklin, 1990; Bain, 2002; Oni et al, 2006; Sickle Cell Society, 2008).

Symptoms
The severity of symptoms varies between patients. Some people have mild symptoms while others are frequently hospitalized for the treatment of more serious complications. However, the basic problem is the same; namely, the sickle-shaped red blood cells tend to get stuck in narrow blood vessels which blocks the flow of blood. This results in the following conditions:

Anaemia
The anaemia in sickle cell disease is caused by red cell destruction or haemolysis. Normally red blood cells live for about 120 days before new ones replace them. Sickle cells usually die after about 10–20 days. The bone marrow cannot make new red blood cells fast enough to replace the dying ones. Anaemia can cause shortness of breath, fatigue, and delayed growth and development in children. The rapid breakdown of red blood cells may also cause yellowing of the eyes and skin which are signs of jaundice (Franklin, 1990; Serjeant and Serjeant, 2001; Sickle cell Society, 2008).

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Key words
- Sickle cell
- Anaemia
- Vaso-occlusive crisis
- Treatment
- Children
- Young people
Clinical

Vaso-occlusive crises (sickle cell crises)

The most common clinical manifestations are painful crises as a result of blockage of small vessels and tissue infarction. These account for more than 90% of all hospital admissions (Platt et al, 1991).

There is enormous variation in the number of crises patients have each year. Some patients have painful crises less than once per year while others have as many as 10 or more episodes (Franklin, 1990; Platt et al, 1991). An average sickle cell crisis usually lasts 4–7 days if there are no complications (Davis et al, 1997). The crisis can come out of nowhere or there can be some precipitating factors, such as dehydration, infection, stress, exposure to extreme temperatures, hypoxia, and strenuous exercise. Repeated crises ultimately result in organ damage and almost any organ can be affected. Other common clinical complications include (Stuart and Nagel, 2004; Sickle Cell Society, 2008):

- Splenic sequestration
- Sepsis, acute chest syndrome
- Priapism (painful erection)
- Lung disease
- Leg ulcers
- Avascular necrosis on the bones
- Kidney damage
- Gallstones

Table 1. Examples of some common types of sickle cell disease

<table>
<thead>
<tr>
<th>Type of Disease</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sickle cell anaemia</td>
<td>Sickle cell anaemia is the most common form of sickle cell disease and it occurs if a person inherits the haemoglobin S gene from both parents. This type tends to be the most serious of the sickle cell conditions.</td>
</tr>
<tr>
<td>Haemoglobin SC disease</td>
<td>Haemoglobin SC disease occurs if a person inherits Hb S from one parent and Hb C from another. On average patients with haemoglobin SC disease have milder symptoms than do those with sickle cell anaemia. Some people with haemoglobin SC disease have a condition that is as severe as any patient with sickle cell anaemia.</td>
</tr>
<tr>
<td>Sickle beta-thalassaemia</td>
<td>Sickle beta-thalassaemia occurs if a person inherits a gene for haemoglobin S from one parent and a gene for beta-thalassaemia from the other.</td>
</tr>
</tbody>
</table>

Figure 1. Examples of blood vessels with normal red blood cells and sickle shaped red blood cells.
Proliferative retinopathy leading to progressive visual loss
Stroke.

Treatment
Sickle cell disease does not have a widely available cure. However, treatments for the symptoms and complications of the disease exist. Bone marrow transplant may offer a cure in a small number of cases. There is some research about gene correction in sickle cell disease (Vichinsky, 2002; Perumbeti and Malik, 2010). Treatment of the complications often includes antibiotics, pain management, intravenous fluids and blood transfusion. Like all patients with chronic diseases they are best managed in a comprehensive multi-disciplinary programme of care.

Prevent infections
In general children and adults with sickle cell anaemia are more vulnerable to infections and have a harder time fighting them off. This is the result of spleen damage from sickled red cells, which prevents it from destroying bacteria in the blood. Giving oral penicillin twice a day beginning at 2 months old can prevent pneumococcal infection and early death. Vaccinations against pneumococcal infections, meningitis, hepatitis and flu are also important (Oni et al, 2006).

Pain management
Pain caused by sickle cell disease can be acute, chronic or a mixture of the two, and patients should be treated accordingly. Acute vaso-occlusive pain is the most common pain in sickle cell disease and is usually treated with a combination of anti-inflammatory agents and opioid or non-opioid analgesics. There are no objective measurements of the severity of pain and analgesia should be titrated against the patient’s reported pain. The severity of pain in sickle cell disease can vary enormously thereby requiring a number of different approaches. Ideally the choice of drug should be influenced by an individual’s analgesic history and some patients may carry cards with details of their ideal analgesic regimen (Okpala and Tawil, 2002; Rees et al, 2003). Paracetamol and non-steroidal anti-inflammatory drugs (NSAIDs) are appropriate for mild to moderate pain. If they are not effective then oral or parenteral opiates can be used. Patients can also benefit from some non-pharmacological approaches such as psychological support, massage, acupuncture, and transcutaneous electrical nerve stimulation.

Distraction and entertainment can be valuable and include television, video games, repeating inspirational phrases, and mental calculations. Studies have suggested that cognitive behavioural therapy for chronic pain can teach patients coping strategies that are also useful for acute pain (Thomas et al, 2001).

It is also important to treat predisposing elements alongside the management of pain. Factors such as diet, poor housing, inadequate heating, lack of education, poor access to expert services, and lifestyle choice may contribute to crises (Westerdale and Jedege, 2004).

Blood transfusion
Although anaemia is a constant feature of sickle cell disease, transfusion is rarely justified for chronic anaemia alone. Transfusion may be required in sickle cell disease either as an emergency measure or to prevent short and long-term complications. Transfusion therapy should be used judiciously because of risks such as:

- Iron overload
- Exposure to hepatitis
- HIV and other infectious agents
- Alloimmunization
- Induction of hyperviscosity
- Limitations on resources.

Transfusion can either be a simple top up or a blood exchange depending on the complications and available resources. The indications for having a blood transfusion

Table 2. Practical advice

Preventing sickle cell crises in school children:

- Make the teachers and other staff aware of sickle cell disease.
- Painful episodes may be prevented by not allowing a child to become over heated or exposed to cold temperatures. A child should always be kept dry and warm.
- Prevent dehydration and encourage plenty of fluid consumption. Let the child have a water bottle with them or allow frequent water breaks. Extra fluid intake and the fact that the kidneys are sometimes affected by sickle cell disease means the child may need bathroom breaks more often than usual. Teachers should be made aware of this.
- Avoid strenuous activities. Anaemia can cause a child with sickle cell disease to tire before others and a rest period may be appropriate. Encourage participation in gym and sports activities but allow a child to stop without attracting undue attention.
- A child should only swim in a heated pool and get dry immediately upon coming out. In some cases swimming should be avoided altogether.
- A child may be absent because of severe pain episodes which can require hospital admission. Support should be given for school work during hospital stays caused by prolonged complications. Children need to go to hospital regularly for follow up.
- Children may have a shorter stature and delayed puberty. The slow rate of growth is caused by a shortage of red blood cells. Taking the vitamin folic acid daily helps to make new red cells.
- Sickle cell disease does not affect intelligence but various problems arising from this lifelong illness may impair academic performance. These should be identified and addressed as they would for any child with difficulties.

Sources: The Sickle Cell Information Centre, 2002; The Learning Trust, City and Hackney Teaching Primary Care Trust, 2006; Oni et al, 2006
Key Points

- The most common types of sickle cell disease are Haemoglobin SS or sickle cell anaemia, Haemoglobin SC disease, and Haemoglobin sickle beta-thalassaemia.
- School nurses need to be well-informed about the condition in order to be able to offer students and their families the best possible support.
- School nurses can help spread awareness of the condition by inviting a speaker from their local sickle cell centre or clinic to educate the entire class and/or the staff about sickle cell disease. They can also become involved in public awareness events.
- Other ways in which they can help include: encouraging blood donations and drives in your community as many sickle cell patients need transfusions, supporting sickle cell research aimed at providing new treatments.

Hydroxyurea

Hydroxyurea is a safe and effective therapy for children and adults with sickle cell anaemia. It mainly works by increasing the level of fetal haemoglobin in the red blood cells thereby reducing the concentration of sickle haemoglobin and sickling itself (Charache et al, 1995; Vichinsky, 2002). Research shows that patients receiving hydroxyurea had a lower incidence of painful events and acute chest syndrome as well as a reduced need for transfusion and hospitalization compared to those receiving no treatment (Charache et al, 1995; Kinney et al, 1999). Hydroxyurea can be used as an alternative to regular blood transfusion. Some practical advice on how to help prevent sickle cell crises in school children can be found in Table 2.

Conclusion

It is important that school nurses have a good understanding of the condition to be able to offer the best possible support and information to the child or young person affected by the condition, peers, parents or carers, and other school employees.

Conflict of interest: None declared

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Further information

Sickle Cell Society

The Society’s mission is to enable and assist individuals with a sickle cell disorder to realise their full economic and social potential. This is achieved by improving opportunities for sickle cell affected individuals and families by raising public awareness through education, advocacy together with the provision of direct welfare services, assisting in research and lobbying.

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Useful resources


Health Workers. Faber and Faber, London