



# Having a simple (top-up) blood transfusion

Information for parents or carers of a child with sickle cell disease (SCD) and thalassaemia



This leaflet will answer some of the questions you have about your child having a simple blood transfusion as a treatment for SCD. If you have any questions or concerns, please speak to the sickle cell doctors or specialist nurses, Snow Fox nurses or the child and adolescent mental health services (CAMHS) practitioner.

## What is blood?

Blood is the liquid in the vessels of the body. It is made up of different types of blood cells and plasma (a strawcoloured liquid that carries the blood cells around the body). The different types of blood cells are:

- red cells, which contain haemoglobin (the red pigment that carries oxygen and carbon dioxide from the tissues of the body)
- white cells (which fight infection)
- platelets (which are clotting cells).

Normally red cells last about 120 days (four months) in the blood (although they last for a shorter time in people with SCD). They are then removed by the body, which means that your body always needs to replace them.

#### What is a blood transfusion?

A blood transfusion is a procedure that puts red cells into the body. This could be a planned procedure, or be performed in an emergency situation.

The two main types of blood transfusions are:

#### A simple transfusion

A few units (bags) of blood are given through a small tube (drip), usually placed in a vein in your child's arm.

#### An exchange blood transfusion

This is a procedure that replaces sickle blood with non-sickle blood (from a donor who does not have SCD). For more information, please speak to the Snow Fox nurses or ask for a copy of the leaflet **Having an exchange blood transfusion:** Information for parents or carers of a child with SCD.

# Why would my child need a simple blood transfusion?

A simple transfusion might be needed if your child's haemoglobin level falls too low. This can happen during a sickle cell crisis, which destroys their red cells. Please contact your GP or hospital doctor immediately if:

- your child's urine is a dark colour
- your child's eyes and skin are yellow (jaundiced)
- your child feels light-headed
- your child feels weak
- your child easily becomes tired or breathless
- your child's spleen feels bigger than normal and child is unwell.

Other reasons for needing a simple transfusion are:

- to protect against longer term sickle cell complications (see below)
- If your child has had abnormal Transcranial Doppler scan results, long-term repeated every 3 to 6 weeks, as a part of the transfusion can help reduce the risk of them having strokes.
- If your child's SCD is very severe, repeated transfusions (every 3 to 6 weeks as part of the longterm transfusion programme) can help reduce the number of crises.
- If your child has had a stroke, long-term repeated transfusions, every 3 to 6 weeks, as a part of the transfusion programme can help reduce the risk of further strokes.
- to get your child ready for an operation to reduce the risk of complications from the general anaesthetic and the surgery

You can find out more from the NHS Blood and Transplant service (contact details at the end of this leaflet).

# **Asking for your consent**

We want to involve you in all the decisions about your child's care and treatment. It is entirely your choice whether they have the transfusion programme (transfusions every 3 to 6 weeks) or not. We will give you relevant information to help you come to a decision with the sickle cell team. It is important that you understand the information and have the time to ask questions and to make your decision.

If you agree to the transfusion for your child, you will be asked to sign a consent form. This confirms that you understand what the treatment involves, and agree for your child to have it. If you would like more information about our consent process, please speak to a member of staff caring for you.

## Is it safe to have a blood transfusion?

In the United Kingdom, we take many precautions to make sure any blood given is as safe as possible. You can find out more from the NHS Blood and Transplant leaflet **Will I need a blood transfusion?** 

The main risk from a transfusion is that the wrong blood could be given by accident. To make sure that the right blood is given, the clinical staff must make careful identification checks. They will ask you to state your child's full name and date of birth and will check the details on their name band, to make sure they receive the correct blood.

They will also regularly monitor your child during the transfusion (check their blood pressure, temperature, pulse and breathing rate) and ask your child how they are feeling.

The risk of contracting a disease, such as hepatitis or human immunodeficiency virus (HIV), is extremely low.

- The risk of getting hepatitis B is less than one in 1.3 million and one in 28 million for hepatitis C.
- The chance of getting HIV from a blood transfusion is one in 6.5 million.

 The possibility of getting variant Creutzfeldt-Jakob disease (vCJD – a rare, incurable brain disease) is extremely small and a number of precautions are taken to reduce this risk.

We now strongly advise that all patients on a transfusion programme are routinely vaccinated against hepatitis B. During the transfusion programme, your child's immunity to hepatitis B and other viral infections will be regularly reviewed.

Although the risk of contracting an infection from blood is very low, we recommend that we check your child for HIV and hepatitis C infection once a year.

# What are the possible complications of blood transfusions?

#### **Minor reactions**

Your child may get a skin rash or a minor fever, for example. These can be treated easily with paracetamol and antihistamines.

#### **Antibodies**

Your child's blood is matched very closely with the blood of the donor (the person who donated the blood). However, it is possible to develop antibodies against the donor blood, so you need to make the clinical staff aware of any symptoms your child may be having after the transfusion (see the section on delayed transfusion reactions). These antibodies can mean that matched blood is harder to find and can take longer to prepare.

#### Iron overload (iron build-up)

This is common in people who receive repeated blood transfusions. When necessary, excess iron can be removed by taking medication (iron chelation) which can be in the form of injections, tablets or liquid. This is much less likely when using the exchange machine, rather than when it is done by hand.

If your child has developed antibodies you will be sent an alert card that says 'I need special blood'. This is to help make sure that anyone treating your child knows that their blood needs to be matched against them. Please make sure you show this to clinical staff before your child receives a transfusion and, in any hospital, where they are treated.

# Why does my child need iron chelation?

Regular blood transfusions can cause a build-up of iron in the body, which over time can cause harm. Our bodies can only remove small amounts of iron and cannot get rid of the build-up of iron from transfusions. Iron chelators are medicines that bind to the excess iron in the blood and help remove it from the body. This stops problems associated with iron build-up. Once the ferritin level (a measure of iron stores) approaches 1,000 you will be offered chelation medicines.

There are three different types of iron chelation medicines. Deferasirox (also known as Exjade®), is a tablet that is taken orally once a day. Deferiprone (also known as Ferriprox®, L1), **is** a tablet or liquid that is taken

orally three times a day. Deferoxamine (also known as Desferal®), is an injectable medicine that is administered over several (usually 8-12) hours, several times a week.

An appropriate leaflet will be provided to you when your child starts iron chelation

## **Delayed transfusion reactions**

Occasionally a patient experiences a delayed transfusion reaction (where the body abnormally breaks down the blood that has been transfused), which may happen within two weeks of the transfusion.

#### This may cause:

- severe generalised sickle cell pain/crisis
- blood in the urine (red or cola colour)
- tiredness
- shortness of breath
- fever
- localised loin/back pain.

If your child experiences these symptoms you must take them to hospital immediately for assessment, and you must inform medical staff that they have been transfused recently.

# Is any preparation needed before the transfusion?

Your child will need to have a blood test before each transfusion. Once the transfusion date is agreed, your child will be given an appointment card from the day unit with the dates for their blood test (this is normally the

day before the transfusion date). If they require 'special blood' (because of antibodies) it is essential that you bring them for blood tests a few days before, or blood may not be available from the transfusion service and their transfusion may be cancelled or delayed.

You can bring your child to the Blood Test Centre on the date on the appointment card, 9am to 5pm. Your child needs to be accompanied by an adult for all tests.

If you are unable to make your appointment, please call Snow Fox Ward or the clinical nurse specialists as soon as possible (contact details at the end of this leaflet).

A blood sample is taken so we know your child's blood group and if they have developed any antibodies. Each time your child has a transfusion, we test their blood against the donated blood, so they are transfused with blood that closely matches their own.

It is important to attend this appointment, so we have time to order the blood that your child needs.

# Where does the transfusion take place?

The transfusion takes place on Snow Fox Ward, first floor, staircase D, South Wing, St Thomas' Hospital unless you are told otherwise. Please arrive before 9am if you have been offered a morning session for the transfusion. If your child has been scheduled for the afternoon session, you must arrive on the ward by 12:30pm.

## Does my child need to bring anything?

Your child might want to bring a book, laptop, or a handheld computer game with them to keep them occupied, as they will need to keep still during the procedure. They should bring their regular daytime medication, including painkillers, as we may not have them on Snow Fox Ward. They should wear loose-fitting clothing.

# What happens during the transfusion?

During a simple blood transfusion, blood is given to your child through a tube (drip), which is usually placed in a vein on their arm. This tube will be removed when the transfusion is finished.

Alternatively, it may be given through a Port-a-Cath (if your child has one). The Port-a-Cath is situated under the skin and uses a large vein in the chest. Cream is normally applied to the port site to numb the area.

The number of units (bags) your child is given depends on how much blood is needed to correct their haemoglobin level. It is usually given in the day unit (9am to 7pm) and your child will usually be given 2 to 3 units of blood. You can both go home once the transfusion is finished. Sometimes, we may ask your child to stay in hospital overnight if it is very late.

# How long will the transfusion take?

The blood transfusion process takes about 2 to 4 hours. This will depend on your child's clinical history and how much blood will be used during the procedure.

# Why does my child need repeat transfusions?

Normal red blood cells only last 120 days in your body. Transfused red cells last less long. Therefore, repeat transfusions are needed to keep the sickle cells in your child's blood at a low level. The frequency depends on the reason for transfusion.

# Can I cancel or change the date of my child's transfusion?

You must contact the sickle cell nurses or Snow Fox Ward as soon as possible so that they can arrange an alternative date (contact details at the end of this leaflet).

## Does it always work?

Most patients selected for a long-term transfusion programme have severe SCD. The top-up transfusion tries to reduce the number of sickle cells in the blood as much as possible, but it can't make them disappear. It also can't suddenly make some of the complications of SCD disappear – like stroke, bone damage or visual loss. This means that sometimes problems can happen to people even though they are on a top-up transfusion programme. Your child will still have an increased risk of infection. It is not a perfect treatment, but it does reduce the chances of new serious problems happening.

### **Further sources of information**

NHS Blood and Transplant service, w: www.nhsbt.nhs.uk/

#### Contact us

Lead consultant for haemoglobin disorders, and consultant in haematology, phone: 020 7188 6203 (secretary), email: Dudhma.Banu@gstt.nhs.uk

**Lead nurse**, **phone**: 020 7188 9432 or 07918 338730 **email**: HaemoglobinopathyCNS@gstt.nhs.uk

**Nurse specialist**, **phone**: 07771 345847 or 07517 592436 **email**: HaemoglobinopathyCNS@gstt.nhs.uk

**Research nurse**, **phone**: 020 7188 6637 or 07517 592436

Specialist nurse – community,

phone: 020 3049 5993 or 07880 781545

Therapy and psychology team, phone: 020 7188 9125 or 07817 905844

Snow Fox Ward

t: 020 7188 4670, Monday to Friday, 7.30am to 8pm.

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