HAVING MY STEM CELL TRANSPLANT

saving the lives of people with blood cancer
HAVING
MY STEM CELL
TRANSPLANT
WHAT’S IN THESE ACTIVITY BOOKLETS

We’ve produced this series of activity booklets for children aged between 5 and 11 who are about to have a stem cell transplant (sometimes also called a bone marrow transplant).

It will help them understand why they need a transplant, what will happen to them and how to look after themselves as they recover. The accompanying guide Helping your child through their stem cell transplant covers the same subjects in more detail.

It’s been written with guidance and advice from parents who have been where you are now, as well as specialist healthcare and teaching professionals, to help you manage your current situation. Each transplant centre will do things differently, so the information in this booklet is a general guide and isn’t intended to replace advice from your child’s doctor and transplant team.

We are here for you and your child on every step of their transplant journey. Our website contains lots of transplant related advice and highlights our other services too.

If you need to ask us any questions, or you would like some more advice, please get in touch with the Anthony Nolan Patient Services team at patientinfo@anthonylogan.org or on 0303 303 0303

PERSONAL DETAILS

This is your activity book. You can write, draw and stick things in it.

It will help you understand more about your stem cell transplant and what happens when you’re in hospital.

My name is

_____________________

I am ______ years old

I live in

_____________________

I am going to be looked after at

_____________________
MEETING YOUR TRANSPLANT TEAM

Now you’ve arrived at hospital, you will be looked after by lots of people. They all have different jobs to do. You might not have met everyone yet, but you will see them as you go.

Here are the hospital staff you’re likely to see. Why not write their names and what they do next to their job? You can draw a picture of them too if you want.

Consultant

Clinical nurse specialists

Nurses

Dietician

Physiotherapist

Hospital teacher

Counsellor or therapist

Porter

Cleaner
Before your transplant, the hospital staff will make sure your body is ready for your new stem cells. This is done using a special kind of medicine called **conditioning therapy**.

For your conditioning therapy you will be given **chemotherapy** and maybe **radiotherapy** as well.

**Chemotherapy** is a medicine the nurse will give you as either a tablet to swallow or through your central line. You will have it every day for up to two weeks. Its job is to remove all the blood cells making you poorly. It also creates space for your new stem cells to live in.

**Radiotherapy** is a type of X-ray. It does the same job as chemotherapy, but you can’t see it or feel it. You will only need it a few times and your doctor might decide you don’t need it at all. It looks like a big machine that will move around you as you lie very still. It’s similar to some of the scans you might have already had.

Unfortunately, conditioning therapy will make you feel poorly. But this means it’s working properly too. The hospital staff will do everything they can to make you feel better and your family will help look after you too.
When our transplant friends had their conditioning therapy, it made them feel poorly too.

**Ava’s chemotherapy** lasted a few days and it made her tummy hurt and she didn’t want to eat her food. She was so tired she stayed in bed lots.

**Tim and Hasan’s chemotherapy** lasted for longer and Tim had to have some radiotherapy too. They were both sick at times and found it hard to eat food. Like Ava, they were both very tired.

The nurses gave them some medicine for their tummies and after a few days they started to feel a bit better.

*How are you feeling after your conditioning therapy?* It’s important to tell the hospital staff if anything is wrong so they can help you.
HAIR LOSS

Your conditioning therapy is likely to make your hair fall out.

It might not happen for a few weeks though. Some children decide to have their hair cut short first to get used to the idea.

Our transplant friends are deciding what they want to do when they lose their hair.

Tim would like to wear a hat to keep his head warm. Can you design some hats for him to try on?

Ava would like to wear a wig. She would like you to help her pick a new one.

Hasan is thinking about having a new colourful haircut. Can you create a new hairstyle for him?

What would you like to do when you lose your hair?

TRANSPLANT DAY

Today is the big day! Today you’re going to get your new stem cells that will soon help you feel better.

They will arrive in a clear plastic bag and look like a red liquid! Your nurse will carefully attach the bag to your central line and the cells will slowly enter your body. It won’t hurt at all. It’s a bit like having a blood transfusion, which you might have had in the past.
FEELING UNWELL AFTERWARDS

Soon after your transplant you will probably feel poorly. It can be caused by your conditioning therapy and because your body has to get used to your new stem cells.

It’s hard to know which parts of your body are going to hurt. Everybody reacts differently and some children feel unwell for longer.

When you do feel unwell, the doctors and nurses will give you medicine or other treatments to help.

Ava was often sick after meals, so the hospital staff changed the types of food she ate. She also had to go to the toilet lots which upset her, but it cleared up when the nurse gave her some medicine.

Tim started to get a red rash on his skin and his eyes were very dry. The doctor told him it was because his body was reacting to his new stem cells. He was given some eye drops and a type of medicine called steroids to stop it happening.

How are you feeling after your transplant?

After his transplant Hasan was very unwell. His whole body ached, and he felt tired all the time. When he felt sick the doctor gave him tablets that made it go away. He also had a sore mouth and gums, so he has a soft toothbrush to clean his teeth.
Lots of children find it very difficult to eat food after their stem cell transplant.

If you're having problems it might be because you feel sick, find it hard to swallow or don't like the taste of some foods.

To make things easier, your doctor might decide you need a feeding tube. This is a bit like your central line, but it carries food into your body, rather than medicine. This isn't food like you normally eat – it’s not mushy vegetables or melted ice cream! It’s all the good bits from food, called nutrients, that you need to feel better.

There are two types of feeding tubes that enter your body at different places.

Some of them go into your nose and down into your tummy. This means you don’t have to worry about your sore throat or how the food tastes.

The other kind go through your skin and into your tummy. You will need to have a small operation, like you did for your central line, to have it fitted. It probably sounds really strange and maybe a little scary to eat your food like this. Please try not to worry though. It’s the best way to make you fit and strong again and you will only need it for a short time.

When you have your tube fitted you will still have to clean your teeth to help keep them healthy.
When you receive your new stem cells it will take a few weeks for them to start making new blood cells.

This means you’re more likely to get an infection from germs that could make you poorly. Normally these germs are removed by your white blood cells.

During this time, you will stay in your own isolation room to keep you safe. It will have a bed, chairs for visitors and a TV. You can have your possessions in there too to make it feel more like home.

The doctors and nurses will check on you often and give you your medicine. They might wear funny looking clothes and masks on their faces so you can’t see their mouth and nose. This is to help stop germs spreading. The room will be cleaned daily to protect you from germs too.

Your grownups can come into the room with you but your brothers and sisters, or other friends might not be allowed to. This is to help stop you getting an infection. They can talk to you on a phone or video call though.
ISOLATION RULES

When you’re in your isolation room there are some rules you and your visitors will have to follow.

Can you design some signs to help everyone remember the following rules and phrases?

Think about other signs you see on roads or in the hospital to help you.

- Visitors must wear protective clothing
- You must stay in your room
- No germs allowed!
- Wash your hands when you arrive and leave
- Only my grownups can come in my room
- Keep everything clean!
THINGS TO DO IN ISOLATION

Staying in isolation might not sound like it’s going to be much fun, but there’s lots of things to do.

Someone called a physiotherapist will come and visit you. They will give you some stretches and gentle exercises to practice. It will make your muscles stronger.

The hospital has its own teacher. They will listen to you read and help you with other school activities in your room if you feel well enough. A play worker will give you some fun activities to do too.

There will be time to watch your favourite films and television programmes. If you have a tablet or games console you will be able to use that too.

If you start to feel sad or upset at any time, there will be someone from the hospital you can talk to.

Your doctor will test your blood everyday to see if your new stem cells are making new blood cells. When they do, you will start to feel much better. This means it will soon be time for you to go home!

You will need to keep looking after yourself as your recovery continues and our next booklet Coming Home After My Stem Cell Transplant will help.
WORD SEARCH

Can you find the following words in the grid opposite? They could be spelt horizontally, vertically or diagonally, forwards or even backwards! You can ask your grownup to help if you like.

Radiotherapy - A type of X-ray which removes the blood cells making you poorly
Chemotherapy - Medicine which removes the blood cells making you poorly
Transplant - When your new stem cells are delivered to your body
Stem cells - Special cells that make other types of blood cells
Isolation - The room you stay in after your transplant
Feeding tube - You might need one if you find it difficult to eat food
Nutrients - The food that goes into your feeding tube
Recovery - The time after your transplant when you start feeling better
Physiotherapist - Gives you exercises to make your body stronger
Nurse - The people who look after you every day in the hospital
Doctor - The person in charge of looking after you
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Going to hospital for my stem cell transplant

Having my stem cell transplant

Coming home after my stem cell transplant

Helping your child through their stem cell transplant

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