

INFORMATION

SHARING THE CARE FOR GOOD NUTRITION

Making the decision to start tube feeding



Where can I find out about tube feeding?

If your child's doctor has asked you to consider tube feeding for your child, you may be experiencing a mixture of emotions - perhaps anxiety, sadness, anger, even guilt. As a parent, of course you want what is best for your child. You want to see that your son or daughter is happy and healthy, but you may also worry that tube feeding in some way will take away the special bond created between you at meal times.

"I had so many questions, practical things, for my child and also for my other kids... I just didn't know what to expect."

You are bound also to have a lot of questions:

How does it work?
How will I cope?
Where do I turn to for support?

Many parents have expressed these same concerns, and have asked for more information at this stage to help them make a more informed decision about tube feeding.



This is not a decision that you have to make alone

Your child's doctor, dietitian or nutrition nurse team should be your first port of call for information and advice about tube feeding. You also may be able to discuss your concerns with a friend, or to ask for advice from another parent or carer who is going through, or has been through, the same experience.

This guide has been developed to help you find out more about tube feeding, give you information on the more practical side of things and suggest other sources of information and support (see the list of associations and organisations on page 11).

"It's the hardest decision I've ever had to make, but it was the right thing to do for my child. Absolutely, I would do it again."



Why has my child's doctor suggested tube feeding?

If your child is experiencing reflux and/or difficulties with chewing and swallowing when eating, your child's doctor will be concerned that he or she might not be getting all the nutrients he or she needs to grow, to play and to learn.

Tube feeds are designed to meet the full nutritional needs of your child, allowing them to get all the nutrients they need. Tube feeding can support your child's growth and development, and in some cases improve their recovery and response to treatments.

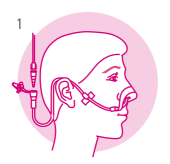
What does tube feeding involve?

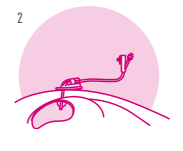
If your child is experiencing feeding difficulties, tube feeding offers an alternative way of feeding that can be adapted to your child's needs.

To prevent the difficulties experienced by your child when taking food by mouth, food in a liquid form is introduced into a tube that goes directly into the digestive system. The ready to feed nutrition, or tube feed, is designed to provide all the essential nutrients and is digested as normal in the gut. Specific formulas are available, if your child has difficulty digesting and/or absorbing nutrients.

There are many types of feeding tubes, NG, NJ, N, PEG. These tubes are shown below feeding via NG/NGT (image 1) or gastrostomy BGT (image 2). You and your child's doctor can discuss together which one is best for your child. There is also more information in our leaflet on the practicalities of tube feeding, which you can obtain from your child's doctor or dietitian.

"I couldn't envisage what it was going to physically look like, once I was shown it, I felt more comfortable...
I was not in the dark anymore."





What do I put in the tube?

The decision about the type of feed that you put into the tube will be taken by you in consultation with the other members of the team that looks after your child's health. Together you will decide what is best for your child.

Ready to feed nutrition made specifically for children who use a feeding tube is ideal. This type of nutrition contains all the required vitamins, minerals, fibres and calories your child needs at the different stages of his or her growth and development, and is designed to flow easily through the tube.

Did you know?

You may be able to give your child his or her medication through the feeding tube - a great benefit if your child has difficulties with swallowing or just doesn't like the taste of the medicine. Your doctor or pharmacist can tell you if your child's medication can be given in this way.

Your child's fluid intake can be controlled using the feeding tube. You can give water or liquid that your child would normally receive orally through the tube to avoid dehydration. It needs to be discussed with the dietitian/doctor if and how much water can be given per day via the feeding tube.



Can my child still take food by mouth once the feeding tube is fitted?

Some parents do continue to offer their child small amounts of normal food to eat in addition to tube feeding.

This not only allows him or her to enjoy the normality of eating different foods together with the family around the table, but also means he or she continues to use and develop the muscles of the mouth (e.g. whilst chewing). Remember, though, to check first with your child's doctor or dietitian in case there are any reasons why he or she should not also be fed in this way.

How will it affect family life?

When making the decision to start tube feeding, one of parents' major concerns is the effect it will have on day-to-day family life. Typical questions include:

Can we still go out to the park?
Can we eat in a restaurant?
Can we go on holiday?
Can we still go to the seaside and swim?

Clearly these are important concerns, as sharing these and other experiences is essential to family bonding and is a key part of growing up and learning to be sociable.

The good news is that all of these activities should be achievable. In fact, the social elements of your family's life definitely can and should continue once the tube is fitted. In many ways, tube feeding can enhance relationships within the family by strengthening existing bonds and providing special moments to share. However, it is a good idea to ask your doctor for advice before undertaking outdoor activities.

"I was really worried that when we were out, people would see Tom's tube and treat him differently... but you can hardly see it."

Who is there to help?

When the decision has been made to start tube feeding, you will have a discussion together with your child's doctor and specialist care/support team (see below) to decide on the best options for your child. The team may include the following healthcare professionals:

Occupational and physical therapists focus on all aspects of your child's movement, including control of the mouth.

A speech-language therapist will assess and help your child's mouth control focusing particularly on speech, language and feeding.

A surgeon may be required to fit your child's feeding tube (but not all tubes require surgery).

The pharmacist
supplies the medication
and nutritional
products that have been
prescribed for your
child and can give you
advice on how it is
to be taken.

Specialist nurses or members of tube feeding associations may carry out home visits to help you and your child with the feeding tube and to provide support and advice.

A dietitian is a specialist in food and nutrition and will be able to give you advice about the best feeding plan for your child.

School nurses may also be involved and will know what happens when the child is at school and potentially feeding, but may also put you in touch with other families who are already tube feeding.

Parent experiences

Remember a practical information brochure is also available, providing you with simple and easy-to-follow instructions and tips at each step of the way. Ask your doctor for more information.



Bonding

"We try and make feeding time as fun and enjoyable as we can. James loves it when we put his favourite songs on. We cuddle and kiss all the time as well, probably even more than before. I am just pleased to know he is putting on weight and getting strong – you can see by his face he is happier."



Trying new foods

"We were really keen for Kate to try new foods even though she was being tube-fed. I think that it is important to give her some taste sensations. She has more confidence now, but to start with she was a bit afraid as she associated it [eating] with being sick. We decided to introduce food in a different way by encouraging her to touch and then smell food and gradually she gained the will to actually try it."

Please note that it is best to ask the doctor or dietitian if it is OK for your child to have foods by mouth before offering them.



Swimming

"Everybody does it their own way, some of the parents I speak to don't like to go out and swim or play in the park as much as before because they worry about the tube. Sam loves swimming so I really wanted him to keep going. Our doctor told us to wait 2 weeks after the tube was fitted and then basically get back into it - so for us it was back to normal, twice a week."

Please note that it is a good idea to seek advice from your child's health care team before undertaking any outdoor activities.

Information, advice and support about **tube feeding**

www.pinnt.com www.cafamily.org.uk www.nutricia.co.uk www.tubefeeding.co.uk For more information please contact your **healthcare professional**.

Reading materials

The following are additional sources of information about tube feeding which you may find helpful:

How do you feel about tube feeding

Practical advice on the signs that may indicate that your child is ready to tube feed.



A practical guide to tube feeding

For a more step by step approach focusing on the more practical aspects of tube feeding.



Glossary of terms

Bolus feeding	A feed that is volume-dependant given over a short period of time	
Clamp	A small device which when pressed closed prevents leakage from the feeding tube	
Continuous feeding	A feed that is given at a pre-set rate, continuously for many hours via pump	
Endoscope	A thin and flexible tube that is passed through the mouth, and down towards the stomach (for investigation and tube placement)	
Enteral feeding	Feed that passes directly into the stomach or intestines	
Extension set	Plastic tubing which can be attached between the feeding set and the feeding tube	
Feeding set	A plastic tube that is attached to the bag with the feed on one end and the tube or extension set on the other	
Flush	Water given to clear the feeding tube/extension tube	
Gastric	Anything related to the stomach	
Gastrostomy	An opening through the abdominal wall into the gastric cavity	
Gastrostomy tube, G-tube	Feeding device through which formula, fluids and/or medication can be given. A G-tube is placed directly into the stomach through an opening in the abdominal wall (stoma) and has balloon end	
Gravity feeding	Feeding that enters the stomach by force of gravity (i.e. placing the feed above the height of the stomach)	
Nasogastric tube, NG-tube	A feeding device through which enteral feed, fluids and/or medication can be given directly into the stomach. An NG-tube is inserted through the nose, passed down the oesophagus and enters directly into the stomach	
Oesophagus	Referred to as the 'swallowing' tube – the path through which food and drink are moved from the mouth to the stomach	
Oral	Anything related to the mouth	
Overnight feeding	Feed that is typically given during the night-time, often by pump	
Pump	A device that delivers feed through the tube at a rate set by you and volume	
Pump feeding	Feeding that is given using a pump	
Stoma	A surgical opening in the abdomen	
Syringe	A medical device used to administer or remove fluid	
Trachea	The 'wind-pipe' that carries air from the mouth to the lungs	
Tube feed, formula	Specifically formulated liquid nutrition that can be tailored to individual needs	
Tube feeding	Supplemental nutrition through a special tube that enters directly into the stomach or intestine	
NJ	An NJ tube is placed usually in hospital and can't be replaced in community. The tube is passed through the nose down the oesophagus and into the small bowel	
Button	A low profile gastrostomy, it is skin level and is retained by a small balloon inside the stomach. These devices are usually changed every 3-6 months at home by a healthcare professional or yourselves if this is something that you want to learn to do	
PEG	A Percutaneous Endoscopic Gastrostomy tube which is placed under general anaesthetic there is a bumper inside to prevent it from coming out. The tube outside the body is approx 20-25cm long	

Your notes			
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four notes			

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- We campaign together for lasting change
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Call our advice line for expert information and advice about caring: **0808 808 7777** (open Monday to Friday, 10am-4pm) **advice@carersuk.org**

We warmly thank all parents and children who contributed to the elaboration of this guide.

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