Information for patients

Surgery for acoustic neuroma

Surgery
Salford Royal NHS Foundation Trust
Stott Lane, Salford, M6 8HD Tel: 0161 206 2303
For further information on this leaflet, its references and sources used, please contact 0161 206 2303
Introduction
This leaflet will offer advice and information to you in preparation for your surgery to remove your acoustic neuroma. It should be read in conjunction with the leaflet “Acoustic Neuroma”.

Preparing for surgery
It is important that you are given as much information as possible to prepare you for your surgery.

Prior to surgery you will be seen in the pre op clinic. Here you will have any investigations needed prior to surgery such as ECG (a tracing of your heart’s electrical activity), blood tests and, if needed, any x-rays that the anaesthetist may request. We ask that you bring along any medication that you are taking.

You will have an opportunity to see the skull base specialist nurse today who will give you information on your surgery and answer any questions you may have. We ask that if you take any medication to thin your blood, i.e., aspirin, warfarin, heparin, clopidogrel, persantin, asasantin, you discuss this with the specialist nurse.

Admission
Day of Surgery
The theatre staff will take you to theatre between 8 and 9 am, though this is dependant on the order of the theatre list. In theatre you will be anaesthetised before entering the main theatre. The anaesthetist will stay with you and monitor you throughout the surgery. Surgery can be lengthy, dependant on the size, location and access required to your tumour and it is not uncommon to be away from the ward for a large part of the day. This does not mean that there is something wrong. The Skull Base surgeons are happy to phone the next of kin or carer when surgery is finished.

What does the surgery entail?
Surgery for acoustic neuroma aims to safely remove the tumour whilst preserving the function of all the important nerves nearby complete tumour removal is usually possible, particularly with small tumours, but not always so. On occasion, your surgeon may decide to leave tiny fragments of tumour if it is felt that complete removal is likely to result in permanent damage of the nearby nerves.

There are 3 main approaches for surgery. Your surgeon will have determined the most appropriate approach for you based
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on your tumour size, location and your hearing function. Your surgeon will shave your hair around where the incision will be made. The length of surgery will depend on the approach taken. Most patients will require a small fat graft which will be taken from your tummy. This will be discussed in more detail in clinic.

Other aspects of Surgery

In all cases the function of the facial nerve is tested with electrical monitoring throughout the operation. There is a risk of facial nerve damage with all approaches of surgery. This can leave you with weakness, loss of movement and inability to close your eye on the affected side. There are differing degrees of function loss dependant on the amount of damage to the nerve. With small tumours the risk of permanent facial weakness is as low as 5%. However, as tumour size increases, the risk to the facial nerve increases such that very large tumours have a risk of permanent facial weakness of around 20%. A higher proportion may have temporary weakness. Apart from removing the tumour, the surgeon’s priority is to maintain facial function and they do all they can to achieve this goal. Occasionally, they may leave a small piece of tumour on the nerve if removing it will put the nerve at higher risk.

Your hearing nerve, the cochlear division of the 8th nerve that your tumour is growing on, is likely to be damaged prior to surgery due to the tumour pressing on this portion of the nerve and the interruption of blood supply. In the majority of people having this surgery there would be little benefit to attempting preservation of the hearing nerve due to this damage and the potential for further damage during surgery.

The translabyrinthine approach will not preserve any hearing and you will wake from the surgery with complete deafness on the affected side. There will be no change to the hearing on the opposite side to the tumour.

You are likely to find yourself imbalanced after this operation due to the loss of function of the vestibular nerve. Most patients compensate well for this and dizziness and imbalance should subside. There are also ways to improve your balance if you find it troubles you.

Other nerves may get damaged during surgery. The 5th cranial nerve, the trigeminal nerve, supplies the sensation to your face. This nerve runs very close to the 8th nerve that the tumour is growing on. It can occasionally get bruised either by the tumour pressing on it or during surgery to remove the tumour. Up to 10% of patients with acoustic neuromas have loss of sensation on the side of the
tumour. As with the other nerve functions, the degree of function loss is dependant on the tumour size and the amount of damage to the nerve. Many patients see a full return of facial sensation with time.

There is a smaller risk to the lower cranial nerves. These supply the swallowing and speech functions of the body. The risk of damage either through tumour compression or damage during surgery is less than 3%. Larger tumours have a higher risk of complications involving these nerves though it must be stressed that this occurs in a very small minority of cases and many patients see a full return of swallowing and voice function.

A complication of surgery is leakage of the fluid surrounding the brain, which occurs in about 3% of patients. This may come through the nose as there is a connection between the nose and the ear, or it may come through the wound. A leak may be treated by draining some of the fluid from around the brain. This is performed by inserting a thin tube into the back. Occasionally it is necessary to close off the leak by performing another smaller operation.

There are several other very rare complications that can occur with this surgery. You will be told about these not because they are common, but because they can be potentially very serious. There is a small chance of infection which can either be superficial and just involve the wound, or can extend internally and affect the linings of the brain. This is meningitis and occurs in about 2% of cases. It is usually treated successfully with intravenous antibiotics. There is a 1% chance of developing a blood clot next to the brain after surgery and this usually needs to be removed with a further operation. There is a small risk (1%) of a stroke from damage either to the blood vessels supplying blood to the brain, or the veins that drain blood from the brain. The severity of such a stroke could be very mild, or very severe and life-threatening. Overall there is a less than 0.5% chance of dying because of the surgery.

Returning to the ward

After a period of time in recovery, you will be transferred to the ward. The ward you are taken to may not be the ward you were admitted to. This may be because you need a higher level of care. Most patients are nursed on ward A3, in the high dependency area or on the main neurosurgical high dependency ward. Occasionally it may be necessary to transfer you to the intensive care unit. This could be due to an existing health issue or a complication during surgery. This does not happen in the majority of cases.
You will have several tubes, lines and monitors attached to you. You may have a tube going into one of the arteries in your wrist to measure your blood pressure constantly. There will also be at least one smaller tube dripping fluids into your veins. You will have leads attached to your chest and arms to measure the activity of your heart and a small probe attached to the end of your finger to measure the levels of oxygen in your body. You will also have either a mask over your nose and mouth or a small tube to the entrance of both nostrils to deliver oxygen to your body.

Most patients have a catheter in place which was inserted in theatre. This will empty your bladder continuously as you will be drowsy for some time after the operation.

You are likely to feel quite sick after the operation and often people are dizzy. You will be given anti-sickness medication, usually via a drip, injection or patch, to help with the sickness. As this improves, you can take oral tablets to reduce the sickness. You may not feel able to tolerate any diet or fluids at this stage, however we do encourage you to at least try and sip small amounts of water. A drip of fluids will be in place to hydrate you.

You may also have some pain where your incision is. The nursing staff will administer pain relief to reduce the pain. Again this is usually via a drip or injection initially.

You will have a bandage around your head and covering the ear on the side of surgery. This applies gentle pressure to prevent any leakage of CSF, the fluid that circulates the brain, and to help the wound to heal. This will stay in place for 2 days. You may also have a wound to your abdomen with a drain coming out of it. It collects any excess of blood to prevent it collecting under the wound. This is where your surgeon took your fat graft from to pack your head wound. This only applies to patients who have had a translabyrinthine approach to surgery.

We expect that you will be very drowsy on this first night. You will be closely monitored overnight and will be woken on an hourly basis to have your blood pressure checked. You will also be asked to answer some questions relating to time and place and the nurse will test your eyes and the power in your arms and legs. This is to check that your brain is functioning well and is a vital part of our monitoring.
Day one

On the first day after your operation you are likely to feel tired, nauseous (sick) and a little dizzy. You may have a mild head pain where the incision was made for your surgery. The nursing staff will be able to administer medication to aid these symptoms. You will be offered breakfast and drinks. We encourage you to try a little though you may not feel like anything at this early stage. Your drip will continue to hydrate you.

The ENT (ears, nose and throat) surgeons and Neurosurgeons will come to see you this morning. You will be asked to make several movements with your face. The doctors are assessing any facial weakness you may have following your operation. Some patients will have some form of weakness at this stage. It does not mean that there has been permanent damage to your facial nerve. It is usually just bruised from the surgery and will take some time to recover. Your surgeon will discuss with you the extent of your facial weakness and the expected outcome. They will also review your wounds, any drainage if you have an abdominal drain in place and ensure that your vital signs, i.e. blood pressure, temperature etc, are within normal limits.

Later in the morning you will have the lines and tubes that have monitored you, removed. You may have to keep the fluid drip in place and are likely to need the catheter for at least another 24 hours until you are more mobile. Your abdominal drain usually comes out this evening. The nurses will assist you with a wash or shower if you feel able to. You should no longer require high dependency care and may be moved to another ward.

It is important that we get you out of bed on the first day to prevent complications such as blood clots and chest infections. You may feel dizzy and tired but the staff will assist you and it will initially be for small periods. You can return to bed any time you wish.

You may have visitors today but we expect that you will feel tired and advise that the visits are short and initially only close relatives or friends.

Day two

You may feel a little brighter today. Try to increase the amount of diet and fluid you are taking. You may still be dizzy and nauseous but you can still have medication to treat that.

The staff and physiotherapists will encourage you to extend the periods you are out of bed. You may want to walk to the bathroom or just up and down the
ward. Ensure you have somebody with you as you can get dizzy quite quickly.

Your bandage will still be in place on your head and you will have a small dressing over your abdominal wound.

The doctors will prescribe an injection into your tummy to help prevent blood clots as you are not as mobile as you would normally be. You will also be prescribed laxatives as we do not want you to strain when going to the toilet as this can raise the pressure of the fluid that cushions the brain and can put you at risk of this fluid leaking from your nose, wound or ear.

If you have facial weakness you may be unable to close your eye fully on the side of the operation. You will be prescribed eye drops to keep the eye wet and may have tape applied to close the eye to prevent it from drying out. You may be referred to an eye specialist if there is very poor closure or signs that the eye is too dry or infected.

Day three, four & five

We expect that you will see some significant improvements within the 3rd, 4th and 5th day. Your nausea should settle and you are likely to feel less dizzy as your body adapts to the loss of the balance nerve on the side of the tumour.

You will feel like eating more and will be able to mobilise a little further each day. You are likely to remain tired for some time so we advise that you do not push yourself too far too soon.

You may find it difficult coming to terms with the changes to your body. If you have lost the hearing on the affected side it can be very frustrating trying to communicate with others. Take your time and ask others to speak slowly and a little louder to help you to hear.

If you have a facial weakness you may feel down at the prospect of a droop on one side of your face. This could occur up to two weeks after the surgery so don’t be alarmed if this develops for the first time after discharge. You can speak to the specialist nurse for more advice. Please be patient with recovery of facial movement as progress can take months. The majority of patients will see significant improvements within 6 months of surgery. Your surgeon will discuss expected outcomes with you after surgery though
recovery is variable.

If you feel wetness from the wound it is important that you inform a member of staff on the ward. This is less likely to happen when you have gone home. Very rarely, patients may get a leak from their nose, down the back of their throat or from the ear. If this occurs please tell a member of staff or if you are at home, you must ring the specialist nurse for advice.

Day six & seven

The majority of patients will go home by day 7. It is important that you have somebody at home with you at least for a day or two. If you do not have anyone who can help you, it may be necessary to refer you to social services for help with your daily activities. You will be discharged with any necessary medications and an information letter from the ward. Your GP will receive a letter informing him of your surgery and any needs you may have. You will only need to see your GP in the event of any complications. If you have any transport needs, please inform the ward staff at the earliest possible moment.

The ward staff will give you a self referral form to see the practice nurse or district nurse to have your clips removed from your head and abdomen if applicable. This is usually on day 10 if the wounds are healthy.

At home

Once you are home we advise you to rest. It is very likely that you will be tired for many weeks and a slow return to normal life is important. It may be useful to take naps in the afternoon and gradually increase the level of activity until your energy returns.

It is equally important that you gently mobilise several times a day to prevent any complications. Take short walks around the house or into the garden and gradually increase the length of your walks.

Do not lift heavy objects or do any strenuous activity for the first 3 months. Continue to take your laxatives as needed to avoid straining.

If you notice any leakage of clear fluid from your nose, down the back of your throat or from the wound, contact the specialist nurse or see your GP.

Driving

The DVLA advises that unless specified by your surgeon, you need not inform them of your surgery unless you are symptomatic of giddiness or, less rarely, seizures. You can clarify this with your surgeon after your operation. We advise that you do not return to driving until fully recovered. This may not apply to those who drive for a living and advice should be sought from the DVLA.
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Working
Dependant on your recovery there is no reason that you should not return to work once fully recovered. This is usually after the first 3 months. Expect to feel a little tired initially. It may be advisable to do a phased return to work, gradually increasing your hours as you feel able. If you have any concerns about returning to work, please discuss this with the surgeons or specialist nurse. You can obtain a certificate of sickness from the ward.

Follow up
The specialist nurse will telephone you after 2 weeks to ensure you are recovering well. If you are unable to use the telephone due to hearing loss, then there is also an email address with the specialist nurse contact or we can liaise with your next of kin or carer with your consent. You will be seen in outpatients at Salford Royal Hospital approximately 6 weeks after your operation. The surgeon will discuss your recovery and can address any issues you may have. Arrangements will be made to follow you up with a scan, usually 1 year after surgery, to ensure there is no sign of residual or reoccurring tumour. This is very unlikely.

There is usually one further scan after 5 years and if this is normal then you will probably be discharged. Follow up scans are carried out as a precaution and it is unlikely you will ever require any further treatment for this tumour.

Hearing aids
If you have lost the hearing on the affected side, you might want to consider a bone anchored hearing aid (BAHA). BAHAs are permanent fixtures surgically inserted into the skull bone behind the ear. There is a detachable sound processor which clips on to the aid and is fairly light. Sound is carried through the bones as your hearing function can no longer use air conduction. BAHAs do not restore your hearing but can improve your everyday hearing function.

CROS Aids consist of 2 hearing aids, one for each ear. The one on the side of the surgery contains a microphone and picks up sounds on this side. These sounds are then sent wirelessly to the other hearing aid in the normal ear where sound is delivered by a tiny speaker. These also help with hearing on the side of surgery but do not replace the hearing you had before. We can discuss either aid if you feel you want to.
Who to contact

If you have any issues after your operation or if you need some help or advice, please contact:

Andrea Wadeson
Skull Base Specialist Nurse
0161 206 2303
07545 513382
Andrea.wadeson@srft.nhs.uk

Jane Riley
Secretary to Mr Rutherford
0161 206 0119

Alison Dapoto
Secretary to Mr King
0161 206 5541

Kim Madden
Secretary to Mr Lloyd and Mr Freeman
0161 206 5754

If the issue is of an urgent nature, see your GP or attend your local accident and emergency department.

Useful contacts

BASIC (Brain and spinal injuries charity)
This is a specialist resource for patients and families affected by neurological diagnosis.
08707500000
Web address: www.basiccharity.org.uk

BANA (British Acoustic Neuroma Association)
This site was developed by people affected by acoustic neuroma and offers information and support to patients and families.
0800 652 3143
Web address: www.bana-uk.com

RNID (Royal National Institute for the Deaf)
This site can offer help and advice to people with hearing difficulties and deafness.
0808 808 0123
Web address: www.rnid.org.uk

BUSK (BAHA users support group)
This site offers advice and support to those considering or who have had a bone anchored hearing aid.
Web address: www.baha-users-support.com
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In accordance with the Equality Act we will make 'reasonable adjustments' to enable individuals with disabilities, to access this treatment / service.

Copies of this information are available in other languages and formats upon request.

Under the Human Tissue Act 2004, consent will not be required from living patients from whom tissue has been taken for diagnosis or testing to use any left over tissue for the following purposes: clinical audit, education or training relating to human health, performance assessment, public health monitoring and quality assurance.

If you object to your tissue being used for any of the above purposes, please inform a member of staff immediately.

Salford Royal operates a smoke-free policy.

For advice on stopping smoking contact the Health and Wellbeing Service on 0161 206 4429

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