As the proportion of older people in our population grows, disorders associated with ageing, such as dementia, Parkinson’s disease and stroke will be even more of a problem. Over 800,000 people in the UK have a diagnosis of dementia right now and this is likely to double in the next 40 years. Scientists still need to understand much more about how different kinds of dementia and other neurodegenerative diseases develop and progress in order to develop better treatments and target them to those who may benefit. Although there have been major advances in medical techniques such as imaging, there is still no real substitute for examining samples of human brain tissue. For many diseases (cancer for instance), tissue removed during surgery can be studied but this is not possible in dementia. Examining brain tissue after someone’s death is still the only accurate way to diagnose the different types of dementia. When looked at together with information about the kinds of symptoms suffered, donated tissue can allow us to see what changes underlie particular memory and neurological problems. In order to do this scientists also need to study tissue from people without disease (‘normal controls’).

We realise that not everyone will wish to consider the idea of donating tissue after death. If you feel it is not for you then we will fully understand if you do not wish to read further.
The Newcastle Brain Tissue Resource (NBTR) is a licensed tissue bank based at Newcastle University. It operates to standards set by the Human Tissue Authority and complies with all current legislation and guidance for working with human tissue. Tissue samples are provided to applicants whose projects are approved by the NBTR committee (this has both scientific and lay members and an independent chair who is not involved in the research). Over the last 40 years research using tissue from the resource has made significant contributions to understanding both normal brain ageing and brain disorders such as: Alzheimer’s disease, Lewy Body disease, vascular dementia, Parkinson's disease, motor neurone disease and depression. Hundreds of scientific papers have been published based on our unique collection of brain tissue. This has contributed to important medical advances such as some of the current drug treatments for Alzheimer’s disease and the identification and treatment of a common type of dementia in older people called Dementia with Lewy Bodies.

**What does participation involve?**

If you decide that you wish to donate your brain for research after your death you would sign a consent form to join our register.

Donations are especially valuable where we have detailed clinical information about the individual and the presence or absence of particular symptoms, so we also ask potential donors to allow us to collect this information. This involves informal assessment interviews with a research nurse, every year for those with a diagnosis of memory impairment, every other year for those without impairment. The assessments are standardised sets of questions and include some questions for a person who knows you well. We also ask whether we may look at your medical records after tissue has been donated. The assessment interviews usually take about two hours and can take place in your own home. Some interviews may take place by telephone for those without a diagnosis of memory impairment. If any areas of concern are identified the research nurse would ask your permission to let your GP know about them.

You might also be asked to give a small blood sample that could be linked later with the donated brain tissue. This is to see whether there may be changes present in the blood during life which might help predict who may later develop dementia or other diseases. If you agreed this would usually involve coming to one of our clinics where trained staff would take a sample of less than 20ml of blood (four teaspoons). As with any blood sampling there is a possible risk of mild discomfort when the sample is taken and that there may be some tenderness or bruising afterwards. If you come to a clinic we will reimburse you for any travelling expenses. This would be purely for research purposes and would not be able to tell
us whether you are at risk of developing problems. It would be completely voluntary and your decision would not affect your participation as a tissue donor. There would be a separate consent form and information sheet for this which you would complete at the time.

When the time came for donation, it would take place in a suitable facility as soon as possible after death. This involves removing and examining the brain and, if possible, the spinal cord and a small blood sample and transporting it to us. The human brain is very complicated and so we usually keep the whole brain (and spinal cord) for diagnosis and research because the location, as well as the nature, of changes may be very important. In some cases we may also ask to examine samples from the heart, blood vessels and nervous system. We would only keep small samples (usually less than 2 cm) of these and specific consent would be sought for this. Although ideally we collect brain and spinal cord, depending on circumstances at the time of donation, it is sometimes only possible to collect the brain.

If you feel that donating brain tissue to NBTR is something you would like to do, you will be given as much information as you require, and opportunities to discuss taking part with the research team and to ask as many questions as you like. You may take as much time as you feel you need to consider your decision. If you decide to take part we will ask you to complete a consent form. If, on the other hand, you decide brain donation is not for you, your decision will be respected and your care will not be influenced in any way.

We are interested in your wishes and feelings about tissue donation even if you do not have legal capacity to consent. If you cannot, or do not wish to, take part in the discussion a consultee (usually a family member or close friend) or nominated representative if you have one, can be asked about your wishes and previously expressed opinions and whether they feel you would have wished to take part in the project. If necessary they can fill in the forms on your behalf.

If in the future you should lose your ability to decide for yourself, it is helpful to have a family member or friend who you trust to uphold your decision to take part. If this should happen without a representative being identified the consent could become invalid. There is a separate information sheet to explain what they need to do.
What would my family have to do when I die?

Normally a family member or representative (person nominated by you to represent you and carry out wishes regarding tissue donation) notifies NBTR when a potential brain donor dies. Your family/representative should contact us as soon as they can to let us know that death has occurred and that they are willing for the brain donation to be carried out. We will keep your family informed of the progress (in accordance with their wishes) and make all the arrangements necessary for the donation to take place. Brain and spinal cord donation does not interfere with, or delay normal funeral arrangements, and does not involve the family in additional expenses. We would provide you and your family or representative with detailed information so they would know exactly what to do when the time comes.

What happens after that?

A very detailed examination of the brain is performed which takes a few months. The research usually begins by examining the tissue using a microscope to find out about any disease processes that have affected the brain. We then use specialist techniques to investigate possible abnormalities in important brain and nerve-cell proteins, and in blood vessels in the brain.

When the examination is complete we usually send a simple letter to your representative/family giving the final diagnosis but the full report is also available if they wish. This report is not written in lay language so it may be more appropriate to send to a G.P. of their choosing, for them to discuss with him/her. If the results have implications for family members this can be addressed at that time.

We also analyse the DNA (genetic material) to look for abnormalities that are specific to some diseases. We will ask for specific consent for this, which you will be at liberty to refuse. Results are not generally in a form that is useful for feedback to individual families. Once diagnosed, tissue would be stored long term in the Newcastle Brain Tissue Resource and be available for use in research into brain disorders.

What kinds of research would my donation be used for?

Many hundreds of scientific papers have already been published using donated tissue from NBTR, in areas such as Alzheimer’s disease, vascular dementia, Lewy Body dementia, frontotemporal dementia, Parkinson’s disease, motor neuron disease and depression. In the past these have contributed to new treatments for Alzheimer’s disease, and recognition of other types of dementia and development of treatments. The samples may be used for research using many kinds of methods and techniques. As knowledge grows, new techniques are being developed all the time.
Many samples are supplied to academic institutions elsewhere within the UK or worldwide. Where samples are sent outside the UK we ask for confirmation that the work will be carried out to the same research and ethical standards.

As it is now possible to grow certain types of cells in the laboratory (stem cells), some researchers want to try to grow these cells using donated tissue. These cells can be extremely useful in that they can be converted into virtually any cell in the body and allow us to understand how different disorders affect cells, tissues, and organs to cause disease. As these cells are potentially capable of growing for very long periods, they will be stored indefinitely. Similar to the access to tissue, access to any cells will be carefully looked at, and the use of any cells produced from donated tissue will be closely regulated.

Sometimes samples are requested by commercial companies such as pharmaceutical companies - often working in partnership with NHS and academic institutions. This is in line with our policy of encouraging work which may lead to future treatments. Neither scientists nor relatives receive any extra financial benefit from this work.

Because human tissue is usually a better alternative, samples are not often used directly in research projects using animals. However, sometimes medical research does involve animals, particularly in the early stages.

You are at liberty to refuse permission for samples to be used for any of these purposes if you wish.

If you or your family would like more information about research supported by NBTR we produce a newsletter, which includes reports on recent studies conducted with donated brain tissue. We will send copies of this on request. If you would like us to do this we will ask permission to store your details for this purpose.

**How is tissue stored?**

We store the tissue in secure, purpose built facilities at Newcastle University to which only authorised staff have access. We freeze some tissue and keep it at a very low temperature so that we can use it for research over a long period (typically 10 years or more). The freezers have alarms and emergency systems so that we can deal quickly with any unexpected rise in temperature and ensure the tissue is always safe. We preserve other tissues in fixative for studies using a microscope.

**Do researchers have to pay for the samples?**

Brain banks are run as ‘not for profit’ organisations but because of the high running costs...
involved, those requesting tissue are asked to make a contribution if possible, using standard tariffs as recommended by the Medical Research Council. However we are flexible and appreciate that sometimes a valuable project may not have sufficient funding for this and would not refuse an application entirely on financial grounds.

What about confidentiality?

To make sure we keep all our information confidential, the samples are issued to researchers labelled only with unique code numbers. We hold records securely on computer and in locked files. The donor’s name is not known to the researchers and you would not be identified in any publications. Once you have signed a consent form your name is placed on a computerised register held at Newcastle University. Only the nurse team who need to keep in touch with you in order to complete the regular assessments and the NBTR manager will have access to this.

How is the tissue disposed of at the end of the research?

Each individual donor may provide samples for many research projects over a period of many years. When we believe we have carried out all possible research on tissue from an individual donor, any remaining material is respectfully disposed of by us, unless other wishes have been stated. Families can withdraw their agreement to use of the tissue at any time, in which case any tissue still held would be disposed of respectfully according to their wishes. Samples already issued cannot be recalled.

What happens next?

We know that discussing tissue donation is a sensitive issue and one about which people have strong feelings. We will totally understand if you do not feel that it is right for you and do not want to proceed. If you do consent you can still withdraw your agreement at any time without having to give a reason. We would recommend that you discuss your decision with your family so that they are aware of your wishes. We are happy to answer any questions or concerns you may have. Our research nurses are also very happy to come and talk things through with family members as well if you would like them to.