What is the purpose of the study?
People with hydrocephalus (fluid in the brain) are often treated with cerebrospinal fluid (CSF) shunts. These are tubes (with a valve) draining the CSF from the fluid cavities in the brain to the abdomen or the heart or around the lungs. These shunts are prone to blockage, over-drainage or infection. In the case of such shunt malfunction, surgery needs to be carried out to rectify it.

Your child has been diagnosed with a shunt problem and needs surgery to make them better. Traditionally we would have replaced part or all of the shunt to treat the problem. This may have been done for your child before. Unfortunately, the revised shunt is also prone to malfunction and nearly half will need further surgery within a year.

More recently, we have been able to employ a different procedure to avoid the need for a shunt altogether. This treatment is called an endoscopic third ventriculostomy (ETV). An ETV involves insertion of a thin telescope (an endoscope) into the ventricles of the brain where the CSF is produced. A hole (ventriculostomy) is made in the floor of one of the ventricles (the third ventricle) to allow the CSF to drain out. A shunt is therefore not needed afterwards. We can remove the shunt or tie it off.
There is evidence to suggest that an ETV can be as good as or maybe better than shunt revision in people that have a blocked shunt. There is definitely a lower risk of infection than with shunts but there is a slightly higher risk of bleeding from the treatment although this risk is small.

The purpose of this study is to determine whether an ETV or shunt revision is a more effective treatment in people with blocked shunts. In order to eliminate any bias, we need to randomly apply either treatment to patients who have a shunt blockage that needs surgery. Overall we need about 450 subjects to reach a valid conclusion and we estimate that the study will take about 5 years.

**How does the study work?**
If you wish your child to partake in the study, we may perform an MRI scan if one has not been done already. This is to determine if ETV is possible. A pre-sealed envelope will then be opened indicating which treatment will be offered. Your child will be treated with either a shunt revision or an ETV. Following surgery, we will keep them in for a few days to make sure the treatment has worked. He/she will be seen about 6 weeks later in the outpatients where further brain imaging will be performed to evaluate the effect of the treatment on the ventricle size.

**What other information will be collected in the study?**
If you wish your child to partake in the study, all their clinical details will be recorded by a neurosurgeon. After the surgery they will be kept in for a few days to make sure they remain well. They will be seen at 6 weeks in clinic and at that stage we will perform an MRI or CT scan to determine how the ventricle size has changed. They will be seen again at 6 months, 1 year, 2 years and 5 years after treatment. We will record whether the treatment worked and whether there were any complications from the treatment.

**Will there be effects on my child's treatment?**
Treatment of your child's condition will continue on in the routine fashion. Participation will not affect him/her in any way. Regardless of whether they enter the study or not and regardless of what type of surgery (ETV or shunt revision) they get, we will treat them the same afterwards and during follow-up as an outpatient.

**Can my child withdraw from the study at any time?**
Yes. You are free to refuse to join the study and may withdraw at any time or choose not to answer certain questions. Your child will receive the same quality of care at the hospital whether you join the study or not.

**Will the information obtained in the study be confidential?**
All data will be kept in the form of hospital registration numbers. Names and addresses of patients will not be used. Even then these hospital numbers can only be accessed by the above named investigators. Anything you say will be treated in confidence, no names will be mentioned in any reports of the study and care will be taken so that individuals cannot be identified from details in reports of the results of the study.

**Will anyone else be told about my participation in the study?**
No
What if I wish to complain about the way in which this study has been conducted?
If you have any cause to complain about any aspect of the way in which you have been approached or treated during the course of this study, the normal National Health Service complaints mechanisms are available to you and are not compromised in any way because you have taken part in a research study.
If you have any complaints or concerns please contact the project coordinator:

Mr. Darach Crimmins,
Leeds General Infirmary
LS1 3EX
(0114) 3928413
darach.crimmins@leedsth.nhs.uk

Otherwise you can use the normal hospital complaints procedure and contact the following person:
Karen Dunwoody
Patient Relations Department,
Ground Floor, Trust Headquarters,
St. James’s University Hospital,
Beckett Street,
Leeds LS9 7TF
(0113) 2066261