



Investigating pain in the developing human brain

Parent Information Leaflet

John Radcliffe Hospital



Oxford University Hospitals
NHS Foundation Trust



Your child is, or may be, eligible to take part in a research study. Before you decide, it is important that you understand why the research is being done and what it involves. Please read the following information carefully and ask us if anything is unclear or if you would like more information.

1. Study title: Investigating pain in the developing human brain

2. What is the purpose of the study?

Infants in hospital often need to have many procedures like blood tests as part of their routine medical treatment, which may cause discomfort. As they cannot tell us how much these procedures hurt, it is difficult to know how much pain they are feeling and to make sure that they receive the right medicines. We know that infants can process discomfort and pain in their brain. By using a special scanner called an MRI (Magnetic Resonance Imaging) scanner, we are able to take detailed 3D pictures of an infant's brain and see how their brain activity changes in response to light, touch, sound or pain. We also know that infants show that they are in pain using different behaviours. These may be indicated by changes in heart rate and breathing in response to pain.

The aim of this research is to understand more about how infants process the outside world and in particular how they feel touch and pain, so that better ways of treating pain can be developed. We are also interested in how infants respond to different stimuli from their environment, such as light and sound, and how this might change across development.

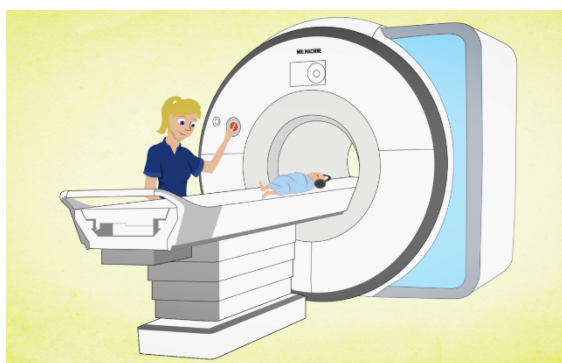
3. Does my child have to take part?

No, it is your decision whether or not your child takes part. If you decide to allow your child to take part, you will be asked to sign a consent form. If you decide you do not want your child to take part, this will not affect your child's care.

If you decide you would like your child to take part, you can change your mind at any time and withdraw your child from the study by telling the research team. You do not have to give a reason. You will be asked if we can use the data/images that have already been collected for analysis (all data) and for publication of results (anonymised data only).

4. What is involved in the study?

If you wish to take part in our study, we will take you and/or your partner and your child to the FMRI Centre, which is where the research MRI scanner is based at the John Radcliffe Hospital. We will use the scanner to record images of your child's brain and its activity. MRI is a safe magnetic imaging technique, which does not involve radiation, and is used to scan infants in hospital. It produces images by measuring changes in blood oxygen levels.



Before the scan, we will check that your child does not have anything metallic on them, and we will give them ear plugs and ear muffs to protect their hearing and to help them sleep soundly through the scan. If possible, giving your child a feed before the scan will make them sleepy and helps to settle them. We will make sure your child is warm and comfortable before we start scanning.

We use a soft touch and a sharp touch device, neither of which pierces the skin but stimulates the receptors we are interested in generally without waking or upsetting the infant. We may also stroke

your child with a soft brush, turn a light on and off, or play a few sounds to them to look at how your child's brain activity changes in response to these stimuli.

During the scan, we may also record your child's heart rate and oxygen saturations using a small probe wrapped around their foot. We may also video your child's face and/or body movements. We may also approach you to ask you if you are happy for us to use these images for teaching, publicity and/or scientific journals. If you agree, we will take separate consent for this as your child's face would be visible in the video footage. This is not a mandatory part of the study. If you choose not to allow us to use the images in this way, this will not affect your child's care or prevent your child from participating in this research.

A clinically trained researcher will be present throughout the study. If your child wakes or needs feeding or changing during the study, we will stop the scanner and ask you whether you wish to continue after their needs have been met. We may ask you to complete a questionnaire following the study.

As we are interested in how your child's response to pain changes as they grow, we may ask if we can study your child more than once during their stay in hospital. We may also ask you if we can contact you in the future, to ask if you would be happy for your child to take part in other research studies. If you agree that we can contact you in the future about other research studies, we will also record your contact details. Your contact details will not be passed onto anyone outside of the research team. You can opt-out of this at any point by contacting Prof Rebecca Slater (details below). Your agreement for us to contact you does not form any obligation to participate in future research.

5. Are there any additional risks or benefits for my child?

A clinically-trained member of the research team will ensure your child's safety at all times. As the MRI scanner is not portable, we need to transport your child to and from the MRI scanning suite in a pram or transport pod. MRI is a safe technique used to image both premature and term infants in hospital. The scanner produces loud repetitive sounds, and we have measured the noise level to ensure that we provide all infants with appropriate ear protection. As the MRI scanner is magnetic, before entering the scanning suite we make sure that neither your child nor any member of the research team is carrying any metal.

Obtaining video footage of your child is non-invasive and does not present any risk to your child. Vital sign monitoring (such as heart rate and breathing rate) has been used clinically for over 20 years without any adverse effects. The sensory stimuli which will be applied to your child have similarly been used in many other patient groups. All studies have a dedicated team of healthcare professionals and researchers that will ensure the safety of your child at all times.

The MRI scan is for research rather than medical diagnosis, so it will not be reviewed by a doctor routinely. If any clinically significant findings are identified at the time of the study then the research team will report these to the clinical care team to handle as appropriate.

There are no direct benefits of participating in this research. This study is designed to gather information to help guide improvements in care for infants in the future. If your child becomes distressed, the research study will be paused or stopped.

6. What information will be collected about my child?

We will collect information about your child from the medical notes, including demographic (e.g. ethnicity), clinical (e.g. number of blood tests in hospital), environmental (e.g. ward transfers) and social factors (e.g. postcode). This information helps us to determine which factors may influence the way an infant copes with pain. In the MRI scanner, we will collect images of your child's brain and its activity, and we may collect information about your child's muscle activity, vital signs, and recordings of their facial expressions. All information and videos that are collected during this research study will be kept strictly confidential. Each infant will be allocated a study number which will be used to label all data.

This study has been registered with the data protection registration office and forms part of an educational programme.

7. What will happen to the results?

Results will be analysed and published in a journal. All publications will be made available on our website <https://neuroimaging.paediatrics.ox.ac.uk>. The findings may also be used for teaching or academic research presentations. No identifying information will be presented about you or your child, unless you have provided specific consent for us to use videos/images of your child in this way.

8. What will happen to my child's data?

We will be using information collected from your child and their medical records in order to conduct this study. Research is a task that we perform in the public interest. The University of Oxford, as Sponsor, is the data controller. This means that we, as University of Oxford researchers, are responsible for looking after the information collected and using it properly. We will use the minimum personally-identifiable information possible. We will keep identifiable information about your child for up to 5 years after the study has finished. This excludes any research documents with personal information, such as consent forms, which will be held securely at the University of Oxford for 25 years after the end of the study.

Data protection regulation provides you with control over your personal data and how it is used. When you agree to your information being used in research, however, some of those rights may be limited in order for the research to be reliable and accurate. Further information about your rights with respect to your personal data is available at <http://www.admin.ox.ac.uk/councilsec/compliance/gdpr/individualrights/>

You can find out more about how we use your information from the contacts in section 12.

Research data may be shared with other researchers doing similar work, both here and abroad. Responsible members of the University of Oxford or the Oxford University Hospitals NHS Trust may be given access to data for monitoring and/or audit of the study to ensure we are complying with regulations.

9. Who is organising and funding this research?

This study is sponsored by University of Oxford and has been funded by The Wellcome Trust. Your doctor will not be paid for including you in this study.

10. Who has reviewed the study?

All research that involves NHS patients has to be approved by a Research Ethics Committee. Approval means that the Committee is satisfied that yours and your child's rights will be respected, that any risks have been reduced to a minimum and balanced against possible benefits, and that

you have been given sufficient information on which to make an informed decision about whether to take part. The South Central Oxford C Research Ethics Committee has reviewed and approved this study

11. Comments or concerns during the study

The University has arrangements in place to provide for harm arising from participation in the study for which the University is the Research Sponsor. NHS indemnity operates in respect of the clinical treatment with which your child is provided. If you wish to complain about any aspect of the way in which you have been approached or treated during the course of this study, you should contact Dr Rebecca Slater (details below) or the University of Oxford Clinical Trials and Research Governance (CTRG) office (01865 (6)16480, ctrg@admin.ox.ac.uk).

12. Contact for further information

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| Prof Rebecca Slater (Study Lead) <i>Professor of Paediatric Neuroscience</i> <i>University of Oxford</i> | 01865 234537 rebecca.slater@paediatrics.ox.ac.uk |
| Dr Eleri Adams (Clinical Lead) <i>Consultant Neonatologist</i> <i>Oxford University Hospitals NHS Trust</i> | 01865 221356 eleri.adams@ouh.nhs.uk |



Picture shows example of an MRI study.

Thank you for reading this information leaflet.