Welcome to the Integrated Research Application System

IRAS Project Filter

The integrated dataset required for your project will be created from the answers you give to the following questions. The system will generate only those questions and sections which (a) apply to your study type and (b) are required by the bodies reviewing your study. Please ensure you answer all the questions before proceeding with your applications.

Please complete the questions in order. If you change the response to a question, please select 'Save' and review all the questions as your change may have affected subsequent questions.

Please enter a short title for this project (maximum 70 characters) The 'Big Baby Trial'				
1. Is your project research?				
Yes No				
2. Select one category from the list below:				
Clinical trial of an investigational medicinal product				
Clinical investigation or other study of a medical device				
Combined trial of an investigational medicinal product and an investigational medical device				
Other clinical trial to study a novel intervention or randomised clinical trial to compare interventions in clinical practice				
Basic science study involving procedures with human participants				
 Study administering questionnaires/interviews for quantitative analysis, or using mixed quantitative/qualitative methodology 				
Study involving qualitative methods only				
 Study limited to working with human tissue samples (or other human biological sample only) 	s) and dat	a (specific project		
Study limited to working with data (specific project only)				
Research tissue bank				
Research database				
If your work does not fit any of these categories, select the option below:				
Other study				
2a. Will the study involve the use of any medical device without a CE Mark, or a CE marked device which has been modified or will be used outside its intended purposes? Yes No				
2b. Please answer the following question(s):				
a) Does the study involve the use of any ionising radiation?	O Yes	No		
b) Will you be taking new human tissue samples (or other human biological samples)?	O Yes	No		
c) Will you be using existing human tissue samples (or other human biological samples)?	O Yes	No		

3. In which countries of the UK will the research sites be located?(Tick all that apply)
☑ England
✓ Scotland
Wales Northern Iroland
Northern Ireland
3a. In which country of the UK will the lead NHS R&D office be located:
● England
Scotland
Wales
○ Northern Ireland
This study does not involve the NHS
4. Which applications do you require?
IRAS Form □ 0 = 5 to 10 to 1
Confidentiality Advisory Group (CAG) Her Majesty's Prison and Probation Service (HMPPS)
Ther majesty same and a robation service (rimin 1.3)
5. Will any research sites in this study be NHS organisations?
Yes No
5a. Are all the research costs and infrastructure costs (funding for the support and facilities needed to carry out research e.g. NHS Support costs) for this study provided by a NIHR Biomedical Research Centre, NIHR Collaboration for Leadership in Health Research and Care (CLAHRC), NIHR Patient Safety Translational Research Centre or Medtech and In Vitro Diagnostic Cooperative in all study sites?
Please see information button for further details.
Please see information button for further details.
5b. Do you wish to make an application for the study to be considered for NIHR Clinical Research Network (CRN) Support and inclusion in the NIHR Clinical Research Network Portfolio?
Please see information button for further details.
The NIHR Clinical Research Network provides researchers with the practical support they need to make clinical studies happen in the NHS e.g. by providing access to the people and facilities needed to carry out research "on the ground".
If you select yes to this question, you must complete a NIHR Clinical Research Network (CRN) Portfolio Application Form (PAF) immediately after completing this project filter question and before submitting other applications. Failing to complete the PAF ahead of other applications e.g. HRA Approval, may mean that you will be unable to access NIHR CRN Support for your study.

6. Do you plan to include any participants who are children?

7. Do you plan at any stage of the project to undertake intrusive research involving adults lacking capacity to consent for themselves?
Answer Yes if you plan to recruit living participants aged 16 or over who lack capacity, or to retain them in the study following loss of capacity. Intrusive research means any research with the living requiring consent in law. This includes use of identifiable tissue samples or personal information, except where application is being made to the Confidentiality Advisory Group to set aside the common law duty of confidentiality in England and Wales. Please consult the guidance notes for further information on the legal frameworks for research involving adults lacking capacity in the UK.
8. Do you plan to include any participants who are prisoners or young offenders in the custody of HM Prison Service or who are offenders supervised by the probation service in England or Wales?
9. Is the study or any part of it being undertaken as an educational project?
10. Will this research be financially supported by the United States Department of Health and Human Services or any of its divisions, agencies or programs?
11. Will identifiable patient data be accessed outside the care team without prior consent at any stage of the project (including identification of potential participants)?

NOTICE OF SUBSTANTIAL AMENDMENT

Please use this form to notify the main REC of substantial amendments to all research other than clinical trials of investigational medicinal products (CTIMPs).

The form should be completed by the Chief Investigator using language comprehensible to a lay person.

Details of Chief Investigator:

Forename/Initials Surname Title

Professor Siobhan Quenby

Clifford Bridge Road, Coventry Work Address

PostCode CV2 2DX

Email s.quenby@warwick.ac.uk

Telephone 07873416716

Fax

For guidance on this section of the form refer to the guidance

Full title of study: Induction of labour for predicted macrosomia

Lead sponsor: University Hospitals Coventry and Warwickshire NHS Trust

Name of REC: South West-Exeter Research Ethics Committee

REC reference number: 18/SW/0039

International Standard Randomised

Controlled Trial Number (ISRCTN):

ISRCTN18229892

ClinicalTrials.gov Identifier (NCT number):

Additional reference number(s):

Ref.Number Description Reference Number

Name of lead R&D office: University Hospitals Coventry & Warwickshire

Date study commenced: 1st Janaury 2018

Protocol reference (if applicable), current

version and date:

V5.0 19th December 2018

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Amendment number and date: AM05 28th February 2019

Гуре of amend	ment
(a) Amendme	nt to information previously given in IRAS
	No No
If yes, plea	ase refer to relevant sections of IRAS in the "summary of changes" below.
(b) Amendme	nt to the protocol
O Yes	No No
	ase submit <u>either</u> the revised protocol with a new version number and date, highlighting changes in document listing the changes and giving both the previous and revised text.
• /	nt to the information sheet(s) and consent form(s) for participants, or to any other supporting n for the study
Yes	○ No
If yes, plea	ase submit all revised documents with new version numbers and dates, highlighting new text in bold.

Is this a modified version of an amendment previously notified and not approved?

Yes

O No

Summary of changes

Briefly summarise the main changes proposed in this amendment. Explain the purpose of the changes and their significance for the study.

If this is a modified amendment, please explain how the modifications address the concerns raised previously by the ethics committee.

If the amendment significantly alters the research design or methodology, or could otherwise affect the scientific value of the study, supporting scientific information should be given (or enclosed separately). Indicate whether or not additional scientific critique has been obtained.

During an interim review of the trial data it has become apparent that the number of women predicted to have a large for gestational age fetus over the 90th centile, that actually deliver a baby over 90th centile is much less than stated in the information sheet. We are therefore cautious of presenting actual figures to participants on the proportion having a baby over the 90th centile. We have therefore amended the participant information sheet to reflect this uncertainty. Any questions the woman might have about the statement that 'women who are told they may have a big baby will not necessarily have a big baby by the time their baby is delivered' will be answered in a personalised discussion with the clinician and in line with usual practice.

Any other relevant information

Applicants may indicate any specific issues relating to the amendment, on which the opinion of a reviewing body is sought.

List of enclosed documents

Document	Version	Date
Participant Information Sheet	6.0	28/02/2019
Cover Letter		

Declaration by Chief Investigator

1. I confirm that the information in this form is accurate to the best of my knowledge and I take full responsibility for it

2. I consider that it would be reasonable for the proposed amendment to be implemented.

This section was signed electronically by Professor Siobhan Quenby on 10/03/2019 18:47.

Job Title/Post: Prof of Obstetrics

Organisation: University of Warwick

Email: s.quenby@warwick.ac.uk

Declaration by the sponsor's representative

I confirm the sponsor's support for this substantial amendment.

This section was signed electronically by Ceri Jones on 07/03/2019 10:13.

Job Title/Post: Head of R&D

Organisation: UHCW NHS Trust

Email: ceri.jones@uhcw.nhs.uk