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) Brain-Computer-Interface to Reduce Fall Risk on Older People		
1. Is your project research?		4
2. Select one category from the list below:	0	·
Clinical trial of an investigational medicinal product	\Diamond	
Clinical investigation or other study of a medical device		
Combined trial of an investigational medicinal product and an investigational medical de	evice	
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 of methodology 	quantitativ	e/qualitative
Study involving qualitative methods only		
Study limited to working with human tissue samples (or other human biological samples) and data (specific project only)		
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:		
if your work does not lit any of those categories, select the option below.		
Other study		
2a. Will the study involve the use of any medical device without a UKCA/CE UKNI/CE Mark, of device which has been modified or will be used outside its intended purposes?	or a UKCA	/CE UKNI/CE marked
2b. Please answer the following question(s):		
a) Does the study involve the use of any ionising radiation?	○ 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 Ireland
3a. In which country of the UK will the lead NHS R&D office be located:
England
Scotland
○ Wales
O Northern Ireland
This study does not involve the NHS
4. Which applications do you require?
▼ IRAS Form
Confidentiality Advisory Group (CAG)
Her Majesty's Prison and Probation Service (HMPPS)
5. Will any research sites in this study be NHS organisations?
5a. Are all the research costs and infrastructure costs (funding for the support and facilities needed to carry out the research e.g. NHS support costs) for this study provided by a NIHR Biomedical Research Centre (BRC), NIHR Applied Research Collaboration (ARC), NIHR Patient Safety Translational Research Centre (PSTRC), or an NIHR Medtech and In Vitro Diagnostic Co-operative (MIC) in all study sites?
Please see information button for further details.
O Von A No
○ Yes No
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 (CRN) provides researchers with the practical support they need to make clinical studies happen in the NHS in England 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, information from your IRAS submission will automatically be shared with the NIHR CRN. Submission of a Portfolio Application Form (PAF) is no longer required.
6. Do you plan to include any participants who are children?

O Yes	No	
7. Do you for themse		trusive research involving adults lacking capacity to consent
○ Yes	No	
loss of cap identifiable Group to s	pacity. Intrusive research means any research wit e tissue samples or personal information, except	or over who lack capacity, or to retain them in the study following the the living requiring consent in law. This includes use of where application is being made to the Confidentiality Advisory in England and Wales. Please consult the guidance notes for volving adults lacking capacity in the UK.
	plan to include any participants who are prisor ffenders supervised by the probation service in	ers or young offenders in the custody of HM Prison Service or n England or Wales?
○ Yes	No	
9. Is the st	tudy or any part of it being undertaken as an ed	lucational project?
Yes	No No	
	is research be financially supported by the Unins, agencies or programs?	ted States Department of Health and Human Services or any of
O Yes	No No	
i e		<u> </u>
	entifiable patient data be accessed outside the identification of potential participants)?	care team without prior consent at any stage of the project
○ Yes	No	
,		

Integrated Research Application System Application Form for

The Chief Investigator should complete this form. The student should complete this form on behalf of the Chief Investigator. Guidance on the questions is available wherever you see this symbol displayed. We recommend reading the guidance first. The complete guidance and a glossary are available by selecting Help.

Please define any terms or acronyms that might not be familiar to lay reviewers of the application.

Short title and version number: (maximum 70 characters - this will be inserted as header on all forms)

PART A: Core study information

1. ADMINISTRATIVE DETAILS

A1. Full title of the research:

A3-1. Chief Investigator:

Title Forename/Initials Surname

Post

Qualifications

ORCID ID

Employer

Work Address

Post Code

Work E-mail

* Personal E-mail

Work Telephone

* Personal Telephone/Mobile

Fax

A copy of a <u>current CV</u> (maximum 2 pages of A4) for the Chief Investigator must be submitted with the application.

A4. Who is the contact on behalf of the sponsor for all correspondence relating to applications for this project? This contact will receive copies of all correspondence from REC and HRA/R&D reviewers that is sent to the CI.

Title Forename/Initials Surname

Address

^{*} This information is optional. It will not be placed in the public domain or disclosed to any other third party without prior consent.

\sim
Post Code
E-mail
Telephone
Fax
A5-1. Research reference numbers. Please give any relevant references for your study:
Applicant's/organisation's own reference number, e.g. R & D (if available):
Sponsor's/protocol number:
Protocol Version:
Protocol Date:
Funder's reference number (enter the reference number or state not applicable):
Project website:
Registry reference number(s): The UK Policy Framework for Health and Social Care Research sets out the principle of making information about research publicly available. Furthermore: Article 19 of the World Medical Association Declaration of Helsinki adopted in 2008 states that "every clinical trial must be registered on a publicly accessible database before recruitment of the first subject"; and the International Committee of Medical Journal Editors (ICMJE) will consider a clinical trial for publication only if it has been registered in an appropriate registry. Please see guidance for more information.
Additional reference number(s):
A5-2. Is this application linked to a previous study or another current application?
○ Yes ○ No
Please give brief details and reference numbers.

2. OVERVIEW OF THE RESEARCH

To provide all the information required by review bodies and research information systems, we ask a number of specific questions. This section invites you to give an overview using language comprehensible to lay reviewers and members of the public. Please read the guidance notes for advice on this section.

A6-1. Summary of the study. Please provide a brief summary of the research (maximum 300 words) using language easily understood by lay reviewers and members of the public. Where the research is reviewed by a REC within the UK Health Departments' Research Ethics Service, this summary will be published on the Health Research Authority (HRA) website following the ethical review. Please refer to the question specific guidance for this question.

A6-2. Summary of main issues. Please summarise the main ethical, legal, or management issues arising from your study and say how you have addressed them.

Not all studies raise significant issues. Some studies may have straightforward ethical or other issues that can be identified and managed routinely. Others may present significant issues requiring further consideration by a REC, R&D office or other review body (as appropriate to the issue). Studies that present a minimal risk to participants may raise complex organisational or legal issues. You should try to consider all the types of issues that the different reviewers may need to consider.

3. PURPOSE AND DESIGN OF THE RESEARCH

A7. Select the appropriate methodology description for this research. Please tick all that apply:
Case series/ case note review
Case control
Cohort observation
☐ Controlled trial without randomisation
☐ Cross-sectional study
☐ Database analysis
☐ Epidemiology
Feasibility/ pilot study
Laboratory study
☐ Metanalysis
Qualitative research
Questionnaire, interview or observation study
Randomised controlled trial
Other (please specify)
A9-2. Is there a sub-study?
○ Yes ○ No ○ Not Answered
A10. What is the principal research question/objective? Please put this in language comprehensible to a lay person.
A11. What are the secondary research questions/objectives if applicable? Please put this in language comprehensible to a lay person.
A12. What is the scientific justification for the research? Please put this in language comprehensible to a lay person.
A13. Please summarise your design and methodology. It should be clear exactly what will happen to the research participant, how many times and in what order. Please complete this section in language comprehensible to the lay person. Do not simply reproduce or refer to the protocol. Further guidance is available in the guidance notes.
A14-1. In which aspects of the research process have you actively involved, or will you involve, patients, service users, and/or their carers, or members of the public?
Design of the research
Management of the research
Undertaking the research
Analysis of results
☐ Dissemination of findings
☐ None of the above

Give details of involvement, or if none please justify the absence of involvement.

A14-2. Have you tested the acceptability of using patient identifiable data in this study without consent?

Please give details.

4. RISKS AND ETHICAL ISSUES

RESEARCH PARTICIPANTS

A15. What is the sample group or cohort to be studied in this research?		
Select all that apply:	67,	
□Blood		
Cancer		
Cardiovascular		
Congenital Disorders		
Dementias and Neurodegenerative [Diseases	
Diabetes		
☐ Ear		
Eye		
Generic Health Relevance		
☐ Infection		
☐ Inflammatory and Immune System		
☐ Injuries and Accidents		
☐ Mental Health		
Metabolic and Endocrine		
Musculoskeletal		
□ Neurological		
Oral and Gastrointestinal		
Paediatrics		
Renal and Urogenital		
Reproductive Health and Childbirth		
Respiratory		
Skin		
Stroke		
Gender:	Male and female participants	
Lower age limit:	Years	
Upper age limit:	Years	

A17-1. Please list the principal inclusion criteria (list the most important, max 5000 characters).

A17-2. Please list the principal exclusion criteria (list the most	important may 5000 characters)

RESEARCH PROCEDURES, RISKS AND BENEFITS

A18. Give details of all non-clinical intervention(s) or procedure(s) that will be received by participants as part of the research protocol. These include seeking consent, interviews, non-clinical observations and use of questionnaires.

Please complete the columns for each intervention/procedure as follows:

- 1. Total number of interventions/procedures to be received by each participant as part of the research protocol.
- 2. If this intervention/procedure would be routinely given to participants as part of their care outside the research, how many of the total would be routine?
- 3. Average time taken per intervention/procedure (minutes, hours or days)
- 4. Details of who will conduct the intervention/procedure, and where it will take place.

A19. Give details of any clinical intervention(s) or procedure(s) to be received by participants as part of the research protocol. These include uses of medicinal products or devices, other medical treatments or assessments, mental health interventions, imaging investigations and taking samples of human biological material. Include procedures which might be received as routine clinical care outside of the research.

Please complete the columns for each intervention/procedure as follows:

- 1. Total number of interventions/procedures to be received by each participant as part of the research protocol.
- 2. If this intervention/procedure would be routinely given to participants as part of their care outside the research, how many of the total would be routine?
- 3. Average time taken per intervention/procedure (minutes, hours or days).
- 4. Details of who will conduct the intervention/procedure, and where it will take place.

A20. Will you withhold an intervention or procedure, which would normally be considered a part of routine care?
A21. How long do you expect each participant to be in the study in total?
A22. What are the potential risks and burdens for research participants and how will you minimise them?
For all studies, describe any potential adverse effects, pain, discomfort, distress, intrusion, inconvenience or changes to lifestyle. Only describe risks or burdens that could occur as a result of participation in the research. Say what steps would be taken to minimise risks and burdens as far as possible.
A23. Will interviews/ questionnaires or group discussions include topics that might be sensitive, embarrassing or upsetting, or is it possible that criminal or other disclosures requiring action could occur during the study?
A24. What is the potential for benefit to research participants?

A25. What arrangements are being made for continued provision of the intervention for participants, if appropriate, once the research has finished? May apply to any clinical intervention, including a drug, medical device, mental health intervention, complementary therapy, physiotherapy, dietary manipulation, lifestyle change, etc.
A26. What are the potential risks for the researchers themselves? (if any)
RECRUITMENT AND INFORMED CONSENT
In this section we ask you to describe the recruitment procedures for the study. Please give separate details for different study groups where appropriate.
A07.4 H
A27-1. How will potential participants, records or samples be identified? Who will carry this out and what resources will be used? For example, identification may involve a disease register, computerised search of social care or GP records, or review of medical records. Indicate whether this will be done by the direct care team or by researchers acting under arrangements with the responsible care organisation(s).
A27-2. Will the identification of potential participants involve reviewing or screening the identifiable personal information of patients, service users or any other person?
∩ Yes ∩ No
Please give details below:
A28. Will any participants be recruited by publicity through posters, leaflets, adverts or websites?
○ Yes ○ No
A29. How and by whom will potential participants first be approached?
A00.4 William abbaic information and the many babally of many and marketing about
A30-1. Will you obtain informed consent from or on behalf of research participants?
○ Yes ○ No
If you plan to seek informed consent from vulnerable groups, say how you will ensure that consent is voluntary and fully informed.
If you are not obtaining consent, please explain why not.
Please enclose a copy of the information sheet(s) and consent form(s).
A30-3. Why is it not practicable for either the researcher's organisation, or the current holder of the information required by the researcher, to seek or obtain patient consent for proposed use of patient identifiable information?
A32. Will you recruit any participants who are involved in current research or have recently been involved in any research prior to recruitment?
○ Yes
○ No
○ Not Known

If Yes, please give details and justify their inclusion. If Not Known, what steps will you take to find out?

A33-1. What arrangements have been made for persons who might not adequately understand verbal explanations or written information given in English, or who have special communication needs?(e.g. translation, use of interpreters)

A34. What arrangements will you make to ensure participants receive any information that becomes available during the course of the research that may be relevant to their continued participation?

A35. What steps would you take if a participant, who has given informed consent, loses capacity to consent during the study? Tick one option only.
The participant and all identifiable data or tissue collected would be withdrawn from the study. Data or tissue which
is not identifiable to the research team may be retained.
The participant would be withdrawn from the study. Identifiable data or tissue already collected with consent would
be retained and used in the study. No further data or tissue would be collected or any other research procedures carried out on or in relation to the participant.
The participant would continue to be included in the study.
Not applicable – informed consent will not be sought from any participants in this research.
O Not applicable – it is not practicable for the research team to monitor capacity and continued capacity will be
assumed.
Further details:

CONFIDENTIALITY

In this section, personal data means any data relating to a participant who could potentially be identified. It includes pseudonymised data capable of being linked to a participant through a unique code number.

A36. Will you be undertaking any of the following activities at any stage (including in the identification of potential participants)?(Tick as appropriate)

Access to medical records by those outside the direct healthcare team

Access to social care records by those outside the direct social care team

Electronic transfer by magnetic or optical media, email or computer networks

Sharing of personal data with other organisations

Export of personal data outside the EEA

Use of personal addresses, postcodes, faxes, emails or telephone numbers

Publication of direct quotations from respondents

Publication of data that might allow identification of individuals

Use of audio/visual recording devices

Storage of personal data on any of the following:

Manual files (includes paper or film)

NHS computers

Social Care Service computers
── Home or other personal computers
University computers
Private company computers
Laptop computers
Further details:
A37. Please describe the physical security arrangements for storage of personal data during the study?
A38. How will you ensure the confidentiality of personal data? Please provide a general statement of the policy and procedures for ensuring confidentiality, e.g. anonymisation or pseudonymisation of data.
Acc Black and for the state of
A39. Please specify whether identifiers will be held in the same database as the clinical data, or in a separate database and linked through a unique study or case number. If held separately, please specify how and at what point the separation will occur. If held in the same database, will the identifiers be encrypted? If so, specify what will be encrypted and who will continue to have access.
A40. Who will have access to participants' personal data during the study? Where access is by individuals outside the direct care team, please justify and say whether consent will be sought.
Storage and use of data after the end of the study
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06-12	2 months
○ 12 mc	onths – 3 years
Over:	3 years
A44. For h	now long will you store research data generated by the study?
Years:	
Months:	
Wioritio.	4
	se give details of the long term arrangements for storage of research data after the study has ended. Say a will be stored, who will have access and the arrangements to ensure security.
Where date	a will be stored, who will have decess and the arrangements to choure security.
INCENTIV	/ES AND PAYMENTS
	research participants receive any payments, reimbursement of expenses or any other benefits or incentives
for taking	part in this research?
	O No
A47. Will in	individual researchers receive any personal payment over and above normal salary, or any other benefits or
	s, for taking part in this research?
○ Yes	○ No
Ores	O NO
	A Y
	s the Chief Investigator or any other investigator/collaborator have any direct personal involvement (e.g. share holding, personal relationship etc.) in the organisations sponsoring or funding the research that may
	to a possible conflict of interest?
- O V	
Yes	O No
	7
NOTIFICA	ATION OF OTHER PROFESSIONALS
	Il you inform the participants' General Practitioners (and/or any other health or care professional responsible are) that they are taking part in the study?
ioi tileli Ca	are, trial tries are taking part in trie study!
Yes	○ No

PUBLICATION AND DISSEMINATION

A50-1. Will the research be registered on a public database?

The UK Policy Framework for Health and Social Care Research sets out the principle of making information about research publicly available. Furthermore: Article 19 of the World Medical Association Declaration of Helsinki adopted in 2008 states that "every clinical trial must be registered on a publicly accessible database before recruitment of the first subject"; and the International Committee of Medical Journal Editors (ICMJE) will consider a clinical trial for publication only if it has been registered in an appropriate registry. Please see guidance for more information.

If Yes, please enclose a copy of the information sheet/letter for the GP/health professional with a version number and date.

Please give details, or justify if not registering the research.			
Please ensure that you have entered registry reference number(s) in q	uestion A5-1.		
A51. How do you intend to report and disseminate the results of the st	tudy?Tick as appropriate:		
Peer reviewed scientific journals			
☐ Internal report			
Conference presentation			
Publication on website			
Other publication	4		
Submission to regulatory authorities			
Access to raw data and right to publish freely by all investigators in	study or by Independent Steering Committee		
on behalf of all investigators			
No plans to report or disseminate the results			
Other (please specify)			
	Cy		
A52. If you will be using identifiable personal data, how will you ensur publishing the results?	e that anonymity will be maintained when		
A53. Will you inform participants of the results?			
○ Yes ○ No	>		
Please give details of how you will inform participants or justify if not do	ina sa		
Please give details of how you will inform participants or justify if not doing so.			
5. Scientific and Statistical Review			
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	?Tick as appropriate:		
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A56. How have the statistical aspects of the research been reviewed Review by independent statistician commissioned by funder or specific of the review by independent statistician Review by company statistician	?Tick as appropriate:		
A56. How have the statistical aspects of the research been reviewed: Review by independent statistician commissioned by funder or specific company independent statistician Review by company statistician Review by a statistician within the Chief Investigator's institution	?Tick as appropriate:		
A56. How have the statistical aspects of the research been reviewed Review by independent statistician commissioned by funder or specific composition. Other review by independent statistician. Review by company statistician. Review by a statistician within the Chief Investigator's institution. Review by a statistician within the research team or multi-centre greater.	?Tick as appropriate:		
A56. How have the statistical aspects of the research been reviewed Review by independent statistician commissioned by funder or specific companies of the review by funder or specific companies of the review by funder or specific companies of the review by independent statistician Review by company statistician Review by a statistician within the Chief Investigator's institution Review by a statistician within the research team or multi-centre graph of the review by educational supervisor	?Tick as appropriate: onsor roup eviewing the statistical aspects. If advice has		
A56. How have the statistical aspects of the research been reviewed: Review by independent statistician commissioned by funder or specific other review by independent statistician Review by company statistician Review by a statistician within the Chief Investigator's institution Review by a statistician within the research team or multi-centre grant Review by educational supervisor Other review by individual with relevant statistical expertise	?Tick as appropriate: onsor roup eviewing the statistical aspects. If advice has		

Work Address
Post Code
Telephone
Fax
Mobile
E-mail
Please enclose a copy of any available comments or reports from a statistician.
A57. What is the primary outcome measure for the study?
A58. What are the secondary outcome measures?(if any)
4
A59. What is the sample size for the research? How many participants/samples/data records do you plan to study in total? If there is more than one group, please give further details below.
Total UK sample size:
Total international sample size (including UK):
Total in European Economic Area:
Further details:
A60. How was the sample size decided upon? If a formal sample size calculation was used, indicate how this was done, giving sufficient information to justify and reproduce the calculation.
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A64-2. Please explain how the responsibilities of sponsorship will be assigned between the co-sponsors listed in A64-1

A65. Has external funding for the research been secured?
Please tick at least one check box.
Funding secured from one or more funders
External funding application to one or more funders in progress
No application for external funding will be made
What type of research project is this?
Standalone project
Project that is part of a programme grant
Project that is part of a Centre grant
Project that is part of a fellowship/ personal award/ research training award
Other
Other – please state:
Outer please state.
A66. Has responsibility for any specific research activities or procedures been delegated to a subcontractor (other than a co-sponsor listed in A64-1)? Please give details of subcontractors if applicable.
○ Yes ○ No
A67. Has this or a similar application been previously rejected by a Research Ethics Committee in the UK or another country?
Yes No
Please provide a copy of the unfavourable opinion letter(s). You should explain in your answer to question A6-2 how the reasons for the unfavourable opinion have been addressed in this application.
A68-1. Give details of the lead NHS R&D contact for this research:
Title Forename/Initials Surname
The Forenane/Initials Suffiance
Organisation
Address
Post Code
Work Email
Telephone
Fax
Mobile
Details can be obtained from the NHS R&D Forum website: http://www.rdforum.nhs.uk

A68-2. Select Local Clinical Research Network for NHS Organisation identified in A68-1:

For more information, please refer to the question specific guidance.	
A69-1. How long do you expect the study to last in the UK?	
Planned start date:	
Planned end date:	
Total duration:	
Years: Months: Days:	
A69-2. How long do you expect the study to last in all countries?	
Planned start date:	
Planned end date:	
Planned end date (clinical interventions):	
Planned end date	
(all trial procedures):	
Total duration:	
Years: Months: Days:	
A71-1. Is this study?	
Single centre	
Multicentre	
A71-2. Where will the research take place? (Tick as appropriate)	
☐ England	
Scotland	
Wales	
Northern Ireland Other countries in European Fearenia Area	
Other countries in European Economic Area	
Does this trial involve countries outside the EU?	
Yes No	
A72. Which organisations in the UK will host the research? Please indicate the type of organisation by ticking the give approximate numbers if known:	box and
NHS organisations in England	
NHS organisations in Wales	
NHS organisations in Scotland	
HSC organisations in Northern Ireland	
GP practices in England	

GP practices in Wales
GP practices in Scotland
GP practices in Northern Ireland
Joint health and social care agencies (eg
community mental health teams)
Local authorities
Phase 1 trial units
Prison establishments
Probation areas
Independent (private or voluntary sector)
organisations
Educational establishments
Independent research units
Other (give details)
Total UK sites in study:
A73-1. Will potential participants be identified through any organisations other than the research sites listed above?
○ Yes ○ No
A74. What arrangements are in place for monitoring and auditing the conduct of the research?
A75-1 What arrangements will be made to review interim safety and efficacy data from the trial? Will a formal data
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Please enclose a copy of relevant documents.	
	and/ or indemnity to meet the potential legal liability of the ing from the design of the research? Please tick box(es) as
through NHS schemes. Indicate if this applies (there is no	ent contracts have designed the research, indemnity is provided oneed to provide documentary evidence). For other protocol please describe the arrangements and provide evidence.
NHS indemnity scheme will apply (protocol authors	with NHS contracts only)
Other insurance or indemnity arrangements will app	ly (give details below)
Please enclose a copy of relevant documents.	
A76-3. What arrangements will be made for insurance a investigators/collaborators arising from harm to partici	
indemnity. Indicate if this applies to the whole study (there	is provided through the NHS schemes or through professional is no need to provide documentary evidence). Where non-NHS practices, please describe the arrangements which will be made at
NHS indemnity scheme or professional indemnity w	ill apply (participants recruited at NHS sites only)
Research includes non-NHS sites (give details of in	surance/ indemnity arrangements for these sites below)
Please enclose a copy of relevant documents.	4
A77. Has the sponsor(s) made arrangements for payme participants where no legal liability arises?	ent of compensation in the event of harm to the research
○ Yes ○ No	
Please enclose a copy of relevant documents.	
A78. Could the research lead to the development of a n	ew product/process or the generation of intellectual property?
○ Yes ○ No ○ Not sure	
Part B: Section 2	
A. General information	A L Y
Information in this sub-section will be included in applicat the research sites.	ions to the Research Ethics Committee and NHS R & D offices at
Is the manufacturer (or other organisation responsiblead sponsor for this study?	le for developing the device) the same organisation named as
Yes No	

Organisation		
Address		
Post Code		
Country		
Telephone		
Fax		
Mobile		
E-mail		
2. Details of the medical devices to be used in the study		
3-1. Further details of the purpose of the study		
Does the study involve:		
Investigation of a new medical device		
Investigation of new implantable material		
Use of an existing product outside the terms of its UKCA/CE UKNI/CE marked intended purpose		
Use of a modified product		
Use of an existing product within its UKCA/CE UKNI/CE marked intended purpose		
3-2. Please give further details below including the following:		
Description of any new device, materials, method of use or operation with a summary of the intended purpose.		
Composition of any new implantable materials, including summary of biocompatibility findings from studies to date.		
A summary of any modifications to UKCA/CE UKNI/CE marked devices.		
A summary of any proposed changes to the UKCA/CE UKNI/CE marked intended purpose.		
For all products with UKCA/CE UKNI/CE mark please attach instructions for use.		

PART C: Overview of research sites

Please enter details of the host organisations (Local Authority, NHS or other) in the UK that will be responsible for the research sites. For further information please refer to guidance.