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# A short introduction to R&D

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# Is this Research?

Different types of projects

- Audit
- Service Development
- Service Evaluation
- Public Health Surveillance
- Case Study
- **Research**

HRA Decision Tool: NRES guidance 'Defining Research'

<http://www.hra-decisiontools.org.uk/research/>

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**NHS**  
Lothian

### Is my study research?

**i** To print your result with title and IRAS Project ID please enter your details below:

Title of your research:

My Research Project

IRAS Project ID (if available):

123-ABC

You selected:

- **'No'** - Are the participants in your study randomised to different groups?
- **'No'** - Does your study protocol demand changing treatment/ patient care from accepted standards for any of the patients involved?
- **'No'** - Are your findings going to be generalisable?

#### Your study would NOT be considered Research by the NHS.

You may still need other approvals.

Researchers requiring further advice (e.g. those not confident with the outcome of this tool) should contact their R&D office or sponsor in the first instance, or the **HRA** to discuss your study. If contacting the HRA for advice, do this by sending an outline of the project (maximum one page), summarising its purpose, methodology, type of participant and planned location as well as a copy of this results page and a summary of the aspects of the decision(s) that you need further advice on to the HRA Queries Line at [HRA.Queries@nhs.net](mailto:HRA.Queries@nhs.net).

For more information please visit the [Defining Research](#) leaflet

[Follow this link to start again.](#)

Print This Page

NOTE: If using Internet Explorer please use browser print function.

# When is NHS ethics required?

- Is it research?
- Does it involve patients?
- Is it legally required to come for NHS review?
- Ethics Favourable Opinion required **before** starting any research study and **before** any recruitment takes place and **before** implement any substantial amendment to the study

# When is R&D approval required?

*Research studies involving NHS patients, their tissues, their data, or NHS resources*

**or**

*Research studies involving NHS staff participating by virtue of their profession*

- ensures the legal obligations of the Board are met
- provides insurance/indemnity for research studies under the Clinical Negligence and Other Risks Indemnity Scheme (CNORIS)
- is a condition of a favourable ethical opinion

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# How to apply for approvals

## IRAS

- web-based application system
- enables you to enter information once about your intended research project - generating all your necessary research approvals
- the only way to apply for approvals (and changes to ongoing studies) in the UK

## Anything else to consider?

- current (short) CV
- GCP certification
- research passports

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# NHS R&D and NRSPCC

NHS Research Scotland Permissions Coordinating Centre (NRSPCC) coordinates the R&D NHS Management approval process and liaises with NHS Board R&D Offices to streamline the process

<http://www.nhsresearchscotland.org.uk>

Generic review – 10 days

(Where a project also involves NHS organisations elsewhere in the UK the study will be supported through existing UK-wide compatibility systems, by which each country accepts the centralised assurances from national coordinating functions)

Local review – 15 days

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# Help!

## General

- Ask ACCORD if you are not sure [accord@nhslothian.scot.nhs.uk](mailto:accord@nhslothian.scot.nhs.uk)
- **It is never too early to talk to us!**

## Ethics

- [www.hra.nhs.uk](http://www.hra.nhs.uk)
- (SESRES) [sesres@nhslothian.scot.nhs.uk](mailto:sesres@nhslothian.scot.nhs.uk)

## IRAS

- Use the information buttons
- Get a project from someone else
- [helpdesk@myresearchproject.org.uk](mailto:helpdesk@myresearchproject.org.uk)

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