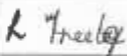


# STANDARD OPERATING PROCEDURE

## S29 – Letters of Access and The Research Passport

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It is the responsibility of all users of this SOP to ensure that the correct version is being used. If you are reading this in a paper format please go [on-line](#) to confirm you have the latest version.

### DISCLAIMER

This generic R&D Standard Operating Procedure (SOP) must be followed unless a study specific SOP exists.

**Once printed this is an uncontrolled document**

Full History			
Version	Date	Author	Reason
1.0		Senior Network R&D Facilitator	New Policy
2.0		SW PenCRN Assistant Manager	Revision to reflect significant changes in NIHR policy
3.0		Senior R&D Facilitator	Updated into Trust Policy template
4		Senior R&D Facilitator	Updated into new Trust Policy template

<b>Associated Trust Policies/ Procedural documents:</b>	<a href="#">S14 Honorary Contracts</a>
<b>Key Words:</b>	<i>Honorary Contracts</i> <i>Letters of Access</i> <i>Research Passport</i>
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1. INTRODUCTION

Research within the NHS can be undertaken by NHS or University / Higher Education Institution (HEI) staff, frequently requiring arrangements for staff to be able to work across a number of NHS organisations.

In accordance with the UK Policy Framework for Health and Social Care Research, UK Health Departments have coordinated the development of The 'HR Good Practice Resource Pack' to help the NHS and other research employers take a consistent approach to handling HR arrangements for those undertaking research in the NHS.

The Royal Devon and Exeter NHS Foundation Trust (hereafter called The Trust) has adopted procedures in line with the above guidance to enable researchers to gain access to the Trust to undertake research-related activity.

2. PURPOSE

This SOP describes the process for issuing Letters of Access (LOA) and outlines when the Research Passport (RP) should be applied for when researchers are undertaking research-related activity in the NHS.

3. SCOPE

This SOP should be read by anyone from a Higher Education Institution (HEI), or employed by another NHS Trust, wishing to undertake research within the Trust.

4. DEFINITIONS

Table with 2 columns: Abbreviation and Full Name. Includes DBS, HEI, HR, HRC, GOG, LOA, R&D, and RP.

5. DUTIES AND RESPONSIBILITIES OF STAFF

Researcher / Supervisor / Trust / University / HEI HR Department

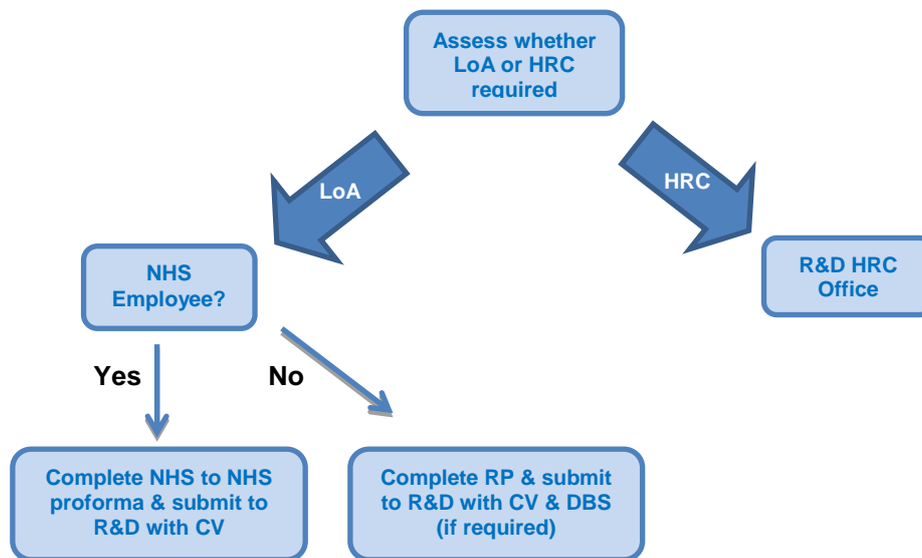
- University / Higher Education Institution (HEI) researchers will be required to complete an RP: relevant sections of the form must be completed by the researcher's Supervisor and University/HEI HR Department OR for those already employed by the NHS, an NHS to NHS Confirmation of Pre-Engagement Checks form must be completed by the researcher's Supervisor.
Human Resources (HR) will determine the level of DBS clearance required and request this (if applicable).
HR will carry out pre-engagement checks including Occupational Health Clearance (if required).
Researcher to produce CV.
Researcher to provide detail to Research and Development (R&D) as to what research related activity they will be involved in.

**R&D Facilitator Team**

- Ensures that a study has received ‘Confirmation of Capacity & Capability’ at the site before issuing a Letter of Access.
- Ensures that all information has been completed correctly on the RP for University/HEI researchers, or NHS to NHS pro forma for NHS researchers.
- Ensure that the relevant pre-engagement checks have been requested by HR to determine whether a LOA or HRC should be issued depending on the nature of the research related activity.

**6. PROCEDURES**

**6.1 Assessment**



R&D will assess whether the researcher requires a LOA or HRC. This is dependent on whether the researcher will have a direct bearing on clinical care. This could be direct contact with patients/service users, children or vulnerable adults and/or has a direct bearing on the quality of care and if access to patient data is required, but only if in NHS facilities. Activities that could have a direct bearing on the quality of care are those that could foreseeably directly affect the type, quality or extent of prevention, diagnosis or treatment of illness, or those that foreseeably cause injury or loss to an individual to whom the organisation has a duty of care.

Researchers with substantive NHS employment contracts or Honorary Clinical Contracts e.g. Clinical Academics with one NHS organisation do not need additional HRCs to conduct research in other NHS organisations. Arrangements for substantive University employees with no NHS HRCs differ depending on whether or not research activities have a direct bearing on the quality of care.

The RP Algorithm (above) can be used to assess when a LoA or HRC is required. LoAs are recommended for researchers working on projects with specified timeframes eg one year and HRCs for longer term studies or researchers working across multiple projects.

For further information on the issuing of HRCs refer to [S14 Honorary Research Contracts](#).

### 6.2 Processing

6.2.1 In order to process a LOA for an NHS employee, the Trust must receive a copy of the following:

- NHS to NHS confirmation of pre-engagement checks form (completed by Supervisor)
- CV (to check employment status and relevant training/qualifications)
- DBS certificate
- Trust to issue LoA

6.2.2 In order to process a LoA for a University/Higher Education Institution, the Trust must receive a copy of the following:

- RP (with relevant sections completed by the institution's HR department and Supervisor)
- CV (to check employment status and relevant training/qualifications)
- DBS certificate
- Trust must complete the relevant section on the RP Form and return a copy to the researcher with their issued LOA.

DBS certificates must be retained for the purpose of processing LoAs only. Original certificates should be returned to the researcher and any copies discarded.

### 6.3 Guidance and template documents for Researchers, R&D and HR Departments

6.3.1 The NIHR [Research in the NHS: Human Resource \(HR\) Good Practice Resource Pack](#) describes the process for handling HR arrangements for researchers and provides a streamlined approach for confirming details of the pre-engagement checks they have undergone with the NHS.

6.3.2 Copies of Trust template letters (NHS and University/HEI templates) and issued LOAs can be found on the shared Drive J:/Research Passports and Letters of Access.

## 7. DISSEMINATION AND TRAINING

7.1 This SOP and associated templates and forms will be uploaded to the [RDE Research website](#) shortly after having been released.

7.2 All staff whose activities are subject to this SOP should ensure that they take time to read and understand the content of this SOP.

7.3 *If applicable, a training log within the Investigator Site File/Trial Master File should be completed to document that members of staff have read and understood the contents of this SOP.*

**8. MONITORING COMPLIANCE AND EFFECTIVENESS OF THIS SOP**

8.1 In order to monitor compliance with this SOP, the auditable standards will be monitored as follows:

No	Minimum Requirements	Evidenced by
1.	R&D section of Research Passport form completed in full.	Check section 8 of Research Passport form
2.	DBS certificates for completed applications should be discarded.	Check that DBS received for application has been deleted from the R&D folder – <a href="#">J:/Research_Develop/ResearchPassports&amp;Letters of Access/IssuedLOAs</a>
3.	Letter of Access workflow completed and relevant documents uploaded to EDGE	Check that a workflow has been created on EDGE for the researcher by searching for the study name 'RD&E HRCs & LOAs'

8.2 Outcomes from audit will be presented to the R&D Governance Oversight Group (GOG) which will monitor any resulting action plans until all issues have been addressed to satisfaction.

8.3 Issues identified via the audit process which require escalation will be referred to GOG.

**9. ARCHIVING ARRANGEMENTS**

9.1 The original of this document will remain with the R&D Quality Assurance Coordinator. An electronic copy will be maintained on the R&D section of the Q-Pulse document management system and a pdf copy on the [RDE Research website](#).

9.2 Archive copies must be maintained for any documents which have been superseded. Archive copies in electronic format should be retained indefinitely.

**10. REFERENCES**

- [Research in the NHS: HR Good Practice Resource Pack](#)
- [UK Policy Framework for Health and Social Care Research](#)