

UHL Implementing Participants in Research Experience Survey (PRES) Research & Innovation SOP C-2033



Trust Reference:
B5/2022

1. Introduction

This Standard Operating Procedure (SOP) describes the procedures to be implemented within the research community when using the National Institute for Health Research (NIHR) Participants in Research Experience Survey (PRES) on UHL sites.

1.0.1)

Every year, the NIHR Clinical Research Network (CRN) asks thousands of participants to give feedback on their experience of taking part in research, as a way of demonstrating to research participants that their contribution is valued, and to help improve the way research studies are designed and delivered, now and in the future.

2. Scope

This SOP applies to all research activity hosted by the University Hospitals of Leicester NHS Trust (UHL) that chooses to implement the NIHR PRES.

3. Version control of PRES

The correct version of the PRES can be found in appendix 1. The PRES is updated annually by the National CRN team and is implemented in line with the new financial year.

3.0.1)

The PRES may NOT be localised. Questions are set nationally and cannot be added to or removed from the document. This is to make sure comparisons can be made between studies, teams, organisations and year-on-year.

4. Distributing the PRES to participants

The PRES should be distributed in paper form in participant study packs with the study name, site and CPMS number (if available) pre-populated by the study team. It is the study team's decision when the most appropriate point in the study would be to share the PRES, but it must be at the same visit for all participants in the study.

4.0.1)

Each study team must reference their study location correctly to support data capture and analysis. All site names must start UHL, followed by the name of the infrastructure. For example, a study in the Biomedical Research Centre must be referenced UHL – BRC. A complete list of codes for research areas is provided in Appendix 3.

4.0.2)

A Perspex box will be provided to research areas on request, where participants can insert completed surveys. It is the study team's responsibility to empty this periodically and send completed surveys to the CRN East Midlands team for input into the database. Participants may wish to take the survey home. In these circumstances freepost envelopes are available from the CRN East Midlands.

4.1)**Digital PRES**

The study team may choose to ask participants to complete the PRES online, provided they have the means to support participants to do so, such as a tablet. The link to the PRES online version is here: <https://crnem.typeform.com/to/WUGMYC?typeform-source=www.leicestershospitals.nhs.uk>

4.2)**Multiple responses from the same participant**

A participant should only receive the PRES once at the allocated study visit, according to the delivery plans of the study team. If participants wish to provide further feedback at later visits, they may scan the QR code on the PRES poster provided as Appendix 2. This poster can be printed, laminated and then populated with the study name, site name and CPMS number using a whiteboard marker.

5. Using the feedback to improve service provision

In agreement with the CRN East Midlands, data from completed surveys will be uploaded into the analysis tool one month from receipt. If negative comments are received, study teams may be contacted by the CRN East Midlands to notify them of the issue to enable the team to respond in real time. Results can be viewed on the live dashboard here:

<https://www.leicestershospitals.nhs.uk/aboutus/education-and-research/leicesters-research/get-involved/pres/>

5.1)

Feedback from the surveys can be used to demonstrate that UHL research listens to its participants and acts on their input to improve the quality of our services. With support from the Research Communications team, study teams can create Patient Engagement boards, 'You Said, We Did' cards to share on social media and in participant correspondence, and use for discussion in team meetings. Information from the PRES can also be used to support disseminating results, in annual reports and in future grant applications.

6. Responsibilities

	Responsibility	Undertaken by	Activity
1	UHL R&I	R&I corporate senior leadership team	Make informed decisions based on patient feedback where it applies to more than one speciality/study type or where an investment is required across the service.
2	UHL R&I	R&I Head of Communications (HoC)	Provide report on PRES responses to R&I Executive and support study teams with participant engagement.
3	CMG/ Specialties	Research Teams	Make informed decisions based on patient feedback where actions can be taken at study/specialty level
4	CRN EM	CRN EM	Upload PRES responses to the dashboard and inform of negative comments in 'real time' to speed up response. Provide information about the breakdown of performance by study, specialty or NIHR infrastructure as requested by HoC
5	R&I Office	Research Manager	Include information in feasibility form, where appropriate, to prompt study teams to consider when they will provide the PRES to their participants.

7. Who Guideline Applies To

This guideline applies to all staff within UHL and external to UHL who are delivering research.

8. Guideline Standards and Procedures

The SOP is detailed so the process can be clearly followed. No flowchart is provided / required.

9. Education and Training

None.

10. Monitoring Compliance

What will be measured to monitor compliance	How will compliance be monitored	Monitoring Lead	Frequency	Reporting arrangements
Sponsor Audit	Randomly chosen for audit	Head of Research Communications	As and when	A report will be produced

11. Supporting Documents and Key References

SOP C-2033 Appendices 1, 2 & 3

12. Key Words

Research, Innovation, Feasibility, PRES, Survey, Feedback, Patient Experience

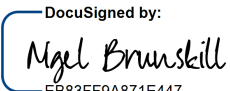
13. Contact and Review Details

CONTACT AND REVIEW DETAILS	
Guideline Lead (Name and Title) Rachael Dowling, Head of Research Communications	Executive Lead Medical director
Details of Changes made during review: Review and update Submit new to PGC	

14.

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This table is used to track the development and approval of the document and any changes made on revised / reviewed versions

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