

Accessible Results: using co-production methods to enable patients with diverse needs to access and understand their blood test results online

Submission date 31/10/2024	Recruitment status Recruiting	☒ Prospectively registered ☒ Protocol
Registration date 04/11/2024	Overall study status Ongoing	☐ Statistical analysis plan ☐ Results
Last Edited 19/08/2026	Condition category Other	☐ Individual participant data ☒ Record updated in last year

Plain English summary of protocol

Background and study aims
NHS England has rolled out online access to test results for patients via the NHS App. Blood tests are the most common type of test requested in primary care. Viewing blood test results online may improve patient engagement and satisfaction. It could also lead to patient anxiety and could mean more patients contact their GP practices to discuss results. Making sure patients can understand their blood results online could therefore have an important impact on primary care workload. Patients need to know what to do next when they receive a blood test result; if this is not communicated well then patients could come to harm. Test result communication is therefore important for patient safety. It is important to make sure that all patients can understand their test results, and that groups such as older people with low digital literacy, minority ethnic groups, and people with disabilities, are not disadvantaged by the move to online blood test result communication. NHS England has acknowledged that the NHS App is not currently fully accessible.
The aim of this study is to develop tools and guidance to help patients with a wide range of needs to access and understand their online blood test results.

Who can participate?

WP1: Patients aged 18 years and over who have had a blood test in the previous month; carers aged 18 years and over of patients who have had a blood test in the previous month; a member of primary care staff involved in systems of blood test communication (to include both clinical and non-clinical staff)
WP2: Patients and carers aged 18 years and over; people aged 18 years and over employed in a role that involves communicating test results to patients, such as GPs, nurses, receptionists, practice managers, healthcare assistants, laboratory staff, clinical biochemists or haematologists.
WP3: Patients (or carers of patients) aged 18 years and over who have had a blood test in the past 1 month; patients (or carers of patients) aged 18 years and over who have been involved in the WP2 workshops

What does the study involve?

This study will use the 'person-based approach', which is an established method for engaging diverse users in developing health interventions. An advisory panel of patients and carers will be involved throughout the three stages of the research (called 'work packages' or WPs). For WP1 the researchers will interview around 40 patients who are having blood tests done in primary care, and 20 primary care staff. They will include patients who have difficulties accessing their results due to disabilities, low digital literacy, or cultural barriers. They will ask them to talk through their test results with us ('think-aloud'). If the patients are not able to view their own results online, the researchers will show them what online blood test results look like, and find out how this could be made more accessible for them. The researchers will ask staff about their attitudes and experiences of the move to online test communication. WP2 will use a series of co-production workshops with patients, carers and healthcare professionals to develop template tools for online blood test results communication. WP3 will test these template tools with a wide range of people, using online surveys. The researchers will use this feedback to develop a final version of the tools. They will share their tools and guidance with stakeholders including the NHS App team, using engagement activities throughout and a dissemination event. This is important to make sure they are used as widely as possible.

What are the possible benefits and risks of participating?

The tools and guidance we develop will be freely available. This work is important to help patients to take ownership of their data and health. Sharing blood test results in a way that is meaningful for patients could also help reduce patient anxiety, improve patient safety and reduce the number of phone calls to GP practices to discuss these results.

Where is the study run from?

University of Bristol (UK)

When is the study starting and how long is it expected to run for?

September 2024 to November 2026

Who is funding the study?

National Institute for Health and Care Research (NIHR) (UK)

Who is the main contact?

Dr Jessica Watson, accessible-results-study@bristol.ac.uk

Contact information

Type(s)

Public, Scientific, Principal investigator

Contact name

Dr Jessica Watson

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Additional identifiers

Integrated Research Application System (IRAS)

349062

Central Portfolio Management System (CPMS)

66092

National Institute for Health and Care Research (NIHR)

159467

Study information

Scientific Title

Accessible Results: enabling patients with diverse needs to access and understand their blood test results online

Study objectives

Improving patient access and understanding of online blood test results is important for patient centred care, patient safety and GP workload.

Ethics approval required

Ethics approval required

Ethics approval(s)

Approved 17/01/2025, North West - Greater Manchester East Research Ethics Committee (3rd Floor, Barlow House 4 Minshull Street, Manchester, M1 3DZ, United Kingdom; +44 (0)207 104 8290, +44 (0)207 104 8061, +44 (0)207 104 8077; gmeast.rec@hra.nhs.uk), ref: 24/NW/0385

Primary study design

Observational

Study design

WP1: qualitative interviews with patients and staff; WP2: co-production workshops to develop draft tools and guidance; WP3: web-based user-testing and refinement of the tools and guidance

Study type(s)

Other

Health condition(s) or problem(s) studied

Promoting the accessibility of online blood test results

Interventions

The person-based approach will be utilised, working with patients, carers, clinicians and other stakeholders. The project will comprise three work packages (WP), building on an ongoing systematic review.

WP1: qualitative interviews with patients and staff to identify user needs.

Six GP practices serving diverse communities in Bristol and Manchester will be recruited, and interviews conducted with around 40 patients and 20 primary care staff. Sampling will include a range of participants, including patients with accessibility needs.

WP2: co-production workshops to develop draft tools and guidance.

8-10 co-production workshops will be conducted involving diverse patient groups, primary care staff and a PPIE advisory panel to develop draft tools and guidance to facilitate the accessibility of online blood test results. These will be user-tested and iteratively refined in subsequent workshops and later WP1 interviews.

WP3: web-based user-testing and refinement of the tools and guidance.

A questionnaire will be disseminated to patients via GP practices and community groups for further user testing and refinement of the tools and guidance.

The final output will be an implementation intervention which may include:

1. Prototype tools for presenting test results in a more accessible way in the NHS App
2. Template wording for clinicians to use when adding comments to test results
3. Posters and/or leaflets for patients to improve their understanding of online test results
4. Resources for patients with diverse needs to help them navigate results online

Intervention Type

Other

Primary outcome(s)

Work package 1: A detailed understanding of patient and staff experiences of online access to test results, collected using qualitative interviews conducted between March - August 2025.

Work package 2: Draft tools and guidance to improve online test communication, developed using co-production workshops involving diverse patient groups, primary care staff and a PPIE advisory panel between May - October 2025

Work package 3: Web-based user-testing and refinement of the tools and guidance using a questionnaire disseminated to patients via GP practices and community groups between Jan - May 2026

Key secondary outcome(s)

There are no secondary outcome measures

Completion date

30/11/2026

Eligibility

Key inclusion criteria

Current key inclusion criteria as of 11/05/2026:

WP1:

Patient and carer interview participants:

Patient aged ≥ 18 years who has had a blood test in the previous month

Or:

Carer aged ≥ 18 of a patient who has had a blood test in the previous month

Primary care staff interview participants:

Member of primary care staff involved in systems of blood test communication (to include both clinical and non-clinical staff)

WP2:

Patient workshops:

Participant aged ≥ 18 years

Or:

Carer aged ≥ 18 years of a patient

Clinician workshops:

Person aged ≥ 18 years employed in a role that involves communicating test results to patients
For example, GPs, nurses, receptionists, practice managers, healthcare assistants, laboratory staff, clinical biochemists or haematologists.

WP3:

Patient (or carer of a patient) aged ≥ 18 years who has had a blood test in the past 6 months

OR

Patient (or carer of a patient) aged ≥ 18 years who has been involved in the WP2 workshops

Previous key inclusion criteria:

WP1:

Patient and carer interview participants:

Patient aged ≥ 18 years who has had a blood test in the previous month

Or:

Carer aged ≥ 18 of a patient who has had a blood test in the previous month

Primary care staff interview participants:

Member of primary care staff involved in systems of blood test communication (to include both clinical and non-clinical staff)

WP2:

Patient workshops:

Participant aged ≥ 18 years

Or:

Carer aged ≥ 18 years of a patient

Clinician workshops:

Person aged ≥ 18 years employed in a role that involves communicating test results to patients
For example, GPs, nurses, receptionists, practice managers, healthcare assistants, laboratory staff, clinical biochemists or haematologists.

WP3:

Patient (or carer of a patient) aged ≥ 18 years who has had a blood test in the past 1 month

OR

Patient (or carer of a patient) aged ≥ 18 years who has been involved in the WP2 workshops

Participant type(s)

Carer, Health professional, Patient

Healthy volunteers allowed

Yes

Age group

Mixed

Lower age limit

18 Years

Upper age limit

100 Years

Sex

All

Total final enrolment

0

Key exclusion criteria

WP1:

Patient and carer interview participants:

1. Clinician feels it would be clinically inappropriate to invite a participant (e.g. patient on palliative care register)
2. Person is unable to give informed consent

Primary care staff interview participants:

Locum staff or staff members working less than 1 day per week in a practice

WP2:

Patient workshops:

1. Person is unable to give informed consent
2. Clinician feels it would be clinically inappropriate to invite a participant (e.g. patient on palliative care register)

WP3:

Clinician feels it would be clinically inappropriate to invite a participant (e.g. patient on palliative care register)

Date of first enrolment

10/02/2025

Date of final enrolment

03/09/2026

Locations**Countries of recruitment**

United Kingdom

England

Study participating centre

NHS Bristol, North Somerset and South Gloucestershire ICB

Floor 2

North Wing

100 Temple Street

Bristol

England

BS1 6AG

Sponsor information

Organisation

University of Bristol

ROR

<https://ror.org/0524sp257>

Funder(s)

Funder type

Government

Funder Name

National Institute for Health and Care Research

Alternative Name(s)

National Institute for Health Research, NIHR Research, NIHRresearch, NIHR - National Institute for Health Research, NIHR (The National Institute for Health and Care Research), NIHR

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be stored in a non-publicly available repository at the University of Bristol Medical School. The raw data will not be shared beyond the core research team. Access to the full anonymised study data set will be limited to members of the immediate research team, the study steering group and authorised representatives from the Sponsor or host institution for monitoring and/or audit of the study to ensure compliance with regulations. Personal data will not be shared with any third parties.

IPD sharing plan summary

Stored in non-publicly available repository

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol file	version 1.3	28/11/2024	13/08/2025	No	No
Protocol file	version 1.4	15/09/2025	09/06/2026	No	No
Study website	Study website	11/11/2025	11/11/2025	No	Yes