

# Improving diversity in clinical research for patients undergoing surgery (PROTECT-DIVERSITY)

<b>Submission date</b> 26/11/2025	<b>Recruitment status</b> Recruiting	<input checked="" type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
<b>Registration date</b> 04/12/2025	<b>Overall study status</b> Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 13/03/2026	<b>Condition category</b> Surgery	<input type="checkbox"/> Individual participant data <input checked="" type="checkbox"/> Record updated in last year

## Plain English summary of protocol

### Background and study aims

The lack of diversity in clinical trials is an important issue, which limits external validity. We are doing this study to check whether consent forms in several different languages can improve access to research for patients having surgery. We will also compare the accuracy and completeness of diversity data collection using different methods such as in your medical records, as part of questionnaires and by asking participants questions.

### Who can participate?

Adults aged 18 years and over undergoing elective surgery.

### What does the study involve?

Patients will be asked to consent to be part of the PROTECT Trial. The consent materials will either be in a traditional paper format or an electronic format, assigned at random. Information such as sex, ethnicity, partnership status, disability, pregnancy status, religion, sexual orientation, gender and gender identity will be collected as part of the PROTECT trial, and patients can refuse to answer any questions they do not want to.

### What are the possible benefits and risks of participating?

The information collected may improve how we collect diversity information for patients taking part in clinical trials in the future. There are no disadvantages to taking part in this study.

### Where is the study run from?

Queen Mary University of London (UK)

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

The study is starting in December 2025 and will run for 2 years.

### Who is funding the study?

The British Journal of Anaesthesia and the Academy of Medical Sciences

Who is the main contact?  
Dr Tom Abbott, protect-admin@qmul.ac.uk

## Contact information

### Type(s)

Scientific, Public, Principal investigator

### Contact name

Dr Tom Abbot

### Contact details

Royal London Hospital  
London  
United Kingdom  
E1 1FR  
+44 203 594 0348  
t.abbott@qmul.ac.uk

## Additional identifiers

### Integrated Research Application System (IRAS)

350756

### Central Portfolio Management System (CPMS)

65329

### Grant Code

WKR0-2023-0016

## Study information

### Scientific Title

Diversity in perioperative research

### Acronym

PROTECT-DIVERSITY

### Study objectives

1. Compare the impact of using multi-lingual consent forms with consent forms in English on the ethnic diversity of patients included in the study.
2. Compare the accuracy and completeness of protected characteristics data collection using different methods.

### Ethics approval required

Ethics approval required

### Ethics approval(s)

approved 10/02/2025, London – South East Research Ethics Committee (2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 207 104 8222; londonsoutheast.rec@hra.nhs.uk), ref: 24/LO/0887

### **Primary study design**

Interventional

### **Allocation**

Randomized controlled trial

### **Masking**

Open (masking not used)

### **Control**

Active

### **Assignment**

Single

### **Purpose**

Health services research

### **Study type(s)**

### **Health condition(s) or problem(s) studied**

Adult patients undergoing elective surgery

### **Interventions**

The master protocol is registered at <https://www.isrctn.com/ISRCTN14639555>

The method of informed consent will use either an electronic or paper method, which is consistent with the master protocol. For this comparison, the mode of consent (electronic or paper) will be determined at random before approaching the patient, which includes consent for entry into the platform. Since the intervention in question is the process of consent, it will not be possible to obtain consent before the 'intervention' i.e. the consent process. In this case, there will be a waiver of consent to randomise to intervention or usual care (electronic consent or paper consent). After randomisation, the patient will complete the consent process using either electronic multi-lingual consent documents or paper consent documents in English, according to group allocation. This process supersedes the procedure for timing of informed consent detailed in the master protocol section nine, which will apply to consent for inclusion in the platform (master protocol) and this comparison. Electronic consent materials will be translated into Polish, Romania, Panjabi, Urdu and Portuguese, which are the five most common languages for people where English/Welsh is not their first language. Translations will be undertaken by an approved/certified provider

### **Intervention Type**

Other

### **Primary outcome(s)**

Reciprocal diversity index for ethnicity on a scale of 0 to 100 at end of trial

### **Key secondary outcome(s)**

1. Completeness of data collection for protected characteristics at trial end
2. Degree of agreement for protected characteristics data between data collection modalities at trial end

### **Completion date**

01/02/2027

## **Eligibility**

### **Key inclusion criteria**

1. Patients aged 18 years and over undergoing elective surgery

### **Participant type(s)**

Patient

### **Healthy volunteers allowed**

No

### **Age group**

Mixed

### **Lower age limit**

18 years

### **Upper age limit**

100 years

### **Sex**

All

### **Total final enrolment**

0

### **Key exclusion criteria**

1. Inability to provide informed consent
2. Co-enrolment in PROTECT CTIMP comparisons
3. Previous enrolment to the PROTECT-DIVERSITY comparison

### **Date of first enrolment**

06/03/2026

### **Date of final enrolment**

01/01/2027

## **Locations**

### **Countries of recruitment**

United Kingdom

England

**Study participating centre**  
**Barts Health NHS Trust**  
The Royal London Hospital  
80 Newark Street  
London  
England  
E1 2ES

## Sponsor information

**Organisation**  
Queen Mary University of London

**ROR**  
<https://ror.org/026zzn846>

## Funder(s)

**Funder type**  
Government

**Funder Name**  
British Journal of Anaesthesia

**Alternative Name(s)**  
British Journal of Anaesthesia Ltd, BJA

**Funding Body Type**  
Private sector organisation

**Funding Body Subtype**  
Other non-profit organizations

**Location**  
United Kingdom

**Funder Name**  
Academy of Medical Sciences

**Alternative Name(s)**

The Academy of Medical Sciences

**Funding Body Type**

Private sector organisation

**Funding Body Subtype**

Universities (academic only)

**Location**

United Kingdom

## Results and Publications

**Individual participant data (IPD) sharing plan****IPD sharing plan summary**

Not expected to be made available

**Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Other files</a>	Consent form	19/06/2025	01/12/2025	No	No
<a href="#">Protocol file</a>	version 3.0	19/06/2025	01/12/2025	No	No