# Study of technologies for the diagnosis of angle closure glaucoma

Submission date 25/01/2023	Recruitment status  No longer recruiting	[X] Prospectively registered		
		[X] Protocol		
Registration date 10/03/2023	Overall study status Completed  Condition category Eye Diseases	Statistical analysis plan		
		Results		
Last Edited		Individual participant data		
05/08/2025		[X] Record updated in last year		

### Plain English summary of protocol

Background and study aims

The ACE study is trying to find out whether people referred by an optician with possible angle closure could be safely diagnosed by healthcare professionals other than eye doctors. Patients will be helping us to determine whether other health professionals besides eye doctors could look after people who have been referred to the eye clinic with angle closure. If the research confirms that this is the case, this will relieve doctors' time in the NHS and doctors could then see patients with serious eye diseases who require treatment more promptly. In the long term, this would potentially help with waiting times in the NHS. If the study shows that having other health professionals see patients once they are stable is not as good as having doctors evaluating them, then this strategy will not be implemented in the NHS.

### Who can participate?

Adults (≥18 years) referred from community optometry to hospital eye services with suspected angle closure

### What does the study involve?

Some images will be obtained from the front of the patient's eyes and the patient will also be seen by an optometrist and an eye doctor at the clinic. Some information about participants will be noted in relation to age, gender, postcode, prescription glasses etc and patients will also be asked to complete a quality-of-life questionnaire.

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

Patients will be helping us to determine whether other health professionals besides eye doctors could look after people referred to the eye clinic with angle closure. If this is the case this will relieve doctors' time in the NHS and doctors could then see patients who need treatment more promptly. Participants may help with waiting times in the NHS. There are no risks associated with the study.

Where is the study run from? Queens University Belfast (UK) When is the study starting and how long is it expected to run for? July 2022 to April 2025

Who is funding the study? National Institute for Health and Care Research (NIHR) Health Technology Assessment Programme (UK)

Who is the main contact:

- 1. Ms Mary Guiney (Public), Ace@nictu.hscni.net
- 2. Prof. Augusto Azura Blanco, (Scientific) a.azuara-blanco@qub.ac.uk

# Contact information

### Type(s)

Principal investigator

#### Contact name

Prof Augusto Azuara-Blanco

### **ORCID ID**

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#### Contact details

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# Type(s)

Public

### Contact name

Mrs Mary Guiney

#### Contact details

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# Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

# Integrated Research Application System (IRAS)

315388

### ClinicalTrials.gov (NCT)

Nil known

### Protocol serial number

B22/08, CPMS 55578

# Study information

#### Scientific Title

Study of technologies for the diagnosis of angle closure glaucoma (ACE)

### Acronym

ACE

### Study objectives

That the two non-contact tests being investigated for diagnosing angle closure glaucoma will be accurate and facilitate a safe and efficient pathway for patients with this condition compared with gonioscopy (reference standard) by an expert consultant ophthalmologist

# Ethics approval required

Old ethics approval format

# Ethics approval(s)

Approved 19/12/2022, London - City & East Research Ethics Committee (Bristol Research Ethics Committee Centre, Whitefriars, Level 3, Block B, Lewins Mead, Bristol, BS1 2NT, UK; +44 (0) 207 104 8171; cityandeast.rec@hra.nhs.uk), ref: 22/LO/0885

# Study design

Prospective cross-sectional multi-centre diagnostic study

# Primary study design

Observational

# Study type(s)

Diagnostic

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

Angle-closure glaucoma

### **Interventions**

This is a diagnostic accuracy study. The tests will be used for triage. The study aims to propose a novel patient pathway. To evaluate the diagnostic performance of two non-contact diagnostic tests compared to gonioscopy (reference standard) by an expert consultant ophthalmologist. Anterior segment optical coherence tomography (AS-OCT) will be interpreted by optometrists

and photographers/imaging technicians and ophthalmologists. Limbal anterior chamber depth (LACD), will be interpreted by optometrists.

There is one visit per patient at the hospital eye clinic services. Testing will be carried out face to face, on an individual basis per patient.

### Intervention Type

Other

### Primary outcome(s)

Sensitivity and specificity of the new pathway to detect angle-closure glaucoma measured using standard formulas and pre-specified criteria for test positivity. All tests will be done at the same clinic visit.

### Key secondary outcome(s))

The following outcome measures will be assessed and recorded in patient notes at the same clinic visit:

- 1. Positive/negative likelihood ratios for angle-closure glaucoma development measured using standard formulas
- 2. Concordance measured using standard formulas
- 3. Long-term health and cost outcomes measured using a Markov model run over an expected lifetime time horizon
- 4. Proportion of patients requiring subsequent clinical assessment by ophthalmologist measured using descriptive statistics
- 5. Proportion of patients unable to undergo tests and of tests of inadequate quality measured using descriptive statistics
- 6. Health-related quality of life measured using the EuroQol Health Questionnaire (EQ-5D-5L)

### Completion date

09/04/2025

# **Eligibility**

### Key inclusion criteria

Adults (≥18 years) referred from community optometry to hospital eye services with suspected angle closure glaucoma

# Participant type(s)

Patient

### Healthy volunteers allowed

No

### Age group

Adult

### Lower age limit

18 years

Sex

### Total final enrolment

641

# Key exclusion criteria

Unable to provide informed consent

### Date of first enrolment

03/04/2023

### Date of final enrolment

30/07/2024

# Locations

### Countries of recruitment

**United Kingdom** 

England

Northern Ireland

Scotland

Wales

# Study participating centre Belfast Health and Social Care Trust

Trust Headquarters A Floor - Belfast City Hospital Lisburn Road Belfast United Kingdom BT9 7AB

# Study participating centre NHS Lothian

Waverley Gate 2-4 Waterloo Place Edinburgh United Kingdom EH1 3EG

# Study participating centre

### Cardiff & Vale University Lhb

Woodland House Maes-y-coed Road Cardiff United Kingdom CF14 4HH

# Study participating centre Moorefields Eye Hospital NHS Foundation Trust

Moorfields Eye Hospital NHS Foundation Trust 162 City Road London United Kingdom EC1V 2PD

# Study participating centre

Nottingham University Hospitals NHS Trust - Queen's Medical Centre Campus

Nottingham University Hospital Derby Road Nottingham United Kingdom NG7 2UH

# Study participating centre Norfolk and Norwich University Hospital

Colney Lane Colney Norwich United Kingdom NR4 7UY

# Study participating centre

Central Manchester University Hospitals NHS Foundation Trust

Trust Headquarters, Cobbett House Manchester Royal Infirmary Oxford Road Manchester United Kingdom M13 9WL

### Study participating centre

# Cambridge University Hospitals NHS Foundation Trust

Cambridge Biomedical Campus Hills Road Cambridge United Kingdom CB2 0QQ

# Study participating centre Sandwell and West Birmingham Hospitals NHS Trust

City Hospital Dudley Road Birmingham United Kingdom B18 7QH

# Study participating centre King's College Hospital NHS Foundation Trust

Denmark Hill London United Kingdom SE5 9RS

# Study participating centre Guy's and St Thomas' NHS Foundation Trust

St Thomas' Hospital Westminster Bridge Road London United Kingdom SE1 7EH

# Study participating centre

# Portsmouth Hospitals University National Health Service Trust

Queen Alexandra Hospital Southwick Hill Road Cosham Portsmouth United Kingdom PO6 3LY

# Study participating centre

# James Paget University Hospitals NHS Foundation Trust

Lowestoft Road Gorleston Great Yarmouth United Kingdom NR31 6LA

# Study participating centre Bedford Hospital NHS Trust

South Wing Kempston Road Bedford United Kingdom MK42 9DJ

# Study participating centre Yeovil District Hospital

Higher Kingston Yeovil United Kingdom BA21 4AT

# Study participating centre York and Scarborough Teaching Hospitals NHS Foundation Trust

York Hospital Wigginton Road York United Kingdom YO31 8HE

# Study participating centre Central Middlesex Hospital NHS Trust

Acton Lane Park Royal London United Kingdom NW10 7NS

# Study participating centre Mid Essex Hospital Broomfield Hospital

Chelmsford United Kingdom CM1 7ET

# Study participating centre Liverpool University Hospitals NHS Foundation Trust

Royal Liverpool University Hospital Prescot Street Liverpool United Kingdom L7 8XP

# Study participating centre University Hospital Southampton NHS Foundation Trust

Southampton General Hospital Tremona Road Southampton United Kingdom SO16 6YD

# Sponsor information

# Organisation

Queen's University Belfast

### **ROR**

https://ror.org/00hswnk62

# Funder(s)

# Funder type

Government

### **Funder Name**

Health Technology Assessment Programme

# Alternative Name(s)

NIHR Health Technology Assessment Programme, Health Technology Assessment (HTA), HTA

# **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 and/or analysed during the current study will be available upon request following the publication of the primary and secondary outcomes. Formal requests for data should be made in writing to Prof. Augusto Azura-Blanco (Chief Investigator) via the Trial Manager, Mary Guiney (ACE@nictu.hscni.net). Requests will be reviewed on a case-by-case basis in collaboration with the Sponsor.

# IPD sharing plan summary

Available on request

### **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<u>Protocol article</u>		04/10/2023	05/10/2023	Yes	No
HRA research summary			26/07/2023	No	No
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes