

# Treatment of severe Diabetic macular oedema with Anti-vascular endothelial growth factor (anti-VEGF) monotherapy versus treatment with anti-VEGF followed by subthreshold Micropulse lasEr when the thickness of the central retina goes below 400 microns: a pragmatic randomised equivalence trial

<b>Submission date</b> 27/09/2024	<b>Recruitment status</b> Recruiting	<input checked="" type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
<b>Registration date</b> 18/11/2024	<b>Overall study status</b> Ongoing	<input type="checkbox"/> Statistical analysis plan <input type="checkbox"/> Results
<b>Last Edited</b> 30/12/2025	<b>Condition category</b> Eye Diseases	<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 macula is the centre of the retina; it gives central sight, colour and fine detail. People with diabetes may develop diabetic macular oedema (DMO). In DMO, fluid leaks from blood vessels and builds up at the macula, causing sight loss. DMO can be mild or severe; this is determined by measuring, in microns ( $\mu\text{m}$ ), how thick the macula is. One  $\mu\text{m}$  is one-thousandth of a millimetre. People presenting with mild DMO (macula less than 400  $\mu\text{m}$  thick; normally it is around 250  $\mu\text{m}$  but varies with sex and ethnicity) are offered macular laser treatment. Laser works well for these patients. Subthreshold micropulse laser (SML), which does not damage the macula, works as well as standard laser, which produces a burn, and is cost-effective.

However, many people present with severe DMO (macula 400  $\mu\text{m}$  or thicker) where the laser does not work well. The standard treatment is eye injections of anti-VEGFs. VEGF stands for vascular endothelial growth factor. VEGF is high in eyes with DMO and causes blood vessel leakage. Anti-VEGFs block VEGF. They are given monthly to begin with, then every 2-3 months for months or years until DMO clears. In many patients DMO comes back after clearing and anti-VEGFs need to be re-started most often monthly initially again.

To improve the care of people with severe DMO this study will compare the current standard care (anti-VEGFs alone) with a strategy in which patients begin with an anti-VEGF but switch to SML once the macula is less than 400  $\mu\text{m}$  thick.

### Who can participate?

Patients aged over 18 years with type 1 or type 2 diabetes and severe DMO

What does the study involve?

Participants are randomly allocated to be treated with either anti-VEGFs alone or anti-VEGFs then SML once the macula is less than 400 µm thick.

What are the possible benefits and risks of participating?

It is considered that the risk associated with the anti-VEGF and SML used within the DAME study is no higher than the risk of standard care. There are a number of expected events associated with the administration of anti-VEGF, SML and intravitreal steroids, Patients will be asked at each visit specifically about each of the following: self-reported central/paracentral scotomas, self-reported reduced colour vision, self-reported metamorphopsia, corneal epithelial erosion, corneal ulcer, endophthalmitis, intraocular inflammation (anterior, posterior or panuveitis), intraocular pressure elevation (over 21 mmHg), intraocular haemorrhage (suprachoroidal /vitreous/pre-retinal haemorrhage), retinal tear, retinal detachment, retinal vasculitis, retinal vascular occlusion (retinal vein or retinal artery occlusion), lens touch (which may occur at the time of an intravitreal injection and may be seen only post-administration in the form of a focal cataract), allergic reaction to any treatments given, including eye drops, angina, myocardial infarction, stroke, transient ischaemic attack (TIA), kidney disease. These events will be collected as safety outcomes and any adverse effects will be monitored by the trial team and DMEC.

Where is the study run from?

Queen's University Belfast (UK)

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

September 2024 to September 2028

Who is funding the study?

National Institute for Health and Care Research (UK)

Who is the main contact?

1. Prof. Noemi Lois, n.lois@qub.ac.uk
2. Mary Guiney, dame@nictu.hscni.net

## Contact information

### Type(s)

Scientific, Principal investigator

### Contact name

Prof Noemi Lois

### Contact details

The Wellcome-Wolfson Institute for Experimental Medicine  
Queen's University Belfast  
Belfast  
United Kingdom  
BT9 7BL  
+44 (0)7484791071  
n.lois@qub.ac.uk

### Type(s)

Public

**Contact name**

Mrs Mary Guiney

**Contact details**

7 Lennoxvale  
Belfast  
United Kingdom  
BT9 5BY  
+44 (0)28 961 51447  
dame@nictu.hscni.net

**Additional identifiers****Clinical Trials Information System (CTIS)**

Nil known

**Integrated Research Application System (IRAS)**

1010626

**Protocol serial number**

24014NL-UC, CPMS 65230

**Study information****Scientific Title**

Treatment of severe Diabetic macular oedema with Anti-vascular endothelial growth factor (anti-VEGF) monotherapy versus treatment with anti-VEGF followed by subthreshold Micropulse lasEr when the thickness of the central retina goes below 400 microns: a pragmatic randomised equivalence trial

**Acronym**

DAME

**Study objectives**

Primary objective:

To determine if the clinical effectiveness of anti-VEGFs and SML is equivalent to anti-VEGF monotherapy

Secondary objectives:

1. To determine the cost-effectiveness of anti-VEGFs and SML compared to anti-VEGF monotherapy via an economic evaluation
2. To evaluate the participant experience and acceptability of anti-VEGFs and SML compared to anti-VEGF monotherapy via a mixed methods evaluation
3. To evaluate the post-trial implementation and scalability of anti-VEGFs and SML via a process evaluation

**Ethics approval required**

Old ethics approval format

## **Ethics approval(s)**

Approved 15/11/2024, South Central - Oxford B Research Ethics Committee (Health Research Authority, 2 Redman Place, Stratford, E20 1JQ, United Kingdom; +44 (0)207 104 8134, +44 (0) 207 104 8019; oxfordb.rec@hra.nhs.uk), ref: 24/SC/0330

## **Study design**

Pragmatic allocation-concealed single-masked (outcome assessors) multicentre randomized (1:1) equivalence trial

## **Primary study design**

Interventional

## **Study type(s)**

Efficacy, Safety

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

Severe diabetic macular oedema (DMO).

## **Interventions**

Comparator Arm: Anti-VEGF Monotherapy (standard care)

Anti-VEGFs including ranibizumab and biosimilars, aflibercept, faricimab, and brolucizumab will be used, as per the standard of care at participating sites. The anti-VEGF should be administered in line with the summary of product characteristics (SmPC).

Intervention Arm: Subthreshold Micropulse Laser (SML)

SML will be applied in line with the DAME Guideline and follow the DAME participant pathway.

Eligible participants who provide consent will be randomised 1:1 to receive SML or to continue with anti-VEGF monotherapy. A minimisation algorithm will be used to ensure balanced allocation of participants across trial arms for potentially important factors including centre, duration of DMO ( $\leq 1$  year,  $> 1$  year), number of doses of anti-VEGFs received up to the time of randomisation (1-6; 7-12), type of anti-VEGF used (ranibizumab, ranibizumab-biosimilar, Brolucizumab, aflibercept, or faricimab) up to the time of randomisation, which will be continued throughout the trial unless lack of efficacy is observed and rescue treatment is needed, presenting BCVA [BCVA  $\geq 69$  ETDRS letters (Snellen equivalent  $\geq 20/40$ ; logMAR  $\geq 0.3$ ), 24–68 ETDRS letters (Snellen equivalent  $\leq 20/50$ -20/320; logMAR 0.4–1.2) and CI-DMO (Yes, No). Minimising randomisation by these variables will ensure both trial arms will be balanced with regard to these potentially important baseline characteristics.

## **Intervention Type**

Drug

## **Phase**

Phase III

## **Drug/device/biological/vaccine name(s)**

Aflibercept, brolucizumab, ranibizumab, faricimab

## **Primary outcome(s)**

Change in best corrected visual acuity (BCVA) in the study eye from randomisation (baseline) to 104 weeks (24 months) (equivalence margin  $\pm 5$  ETDRS letters)

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

All measured at 104 weeks (24 months) from randomisation:

1. Central Retinal Thickness in the study eye. CRT in the central 1 mm of the retina as measured using Spectral-Domain optical Coherence Tomography (SD- OCT)
2. Health-related and vision-related quality of life. National Eye Institute Visual Function Questionnaire (NEI VFQ) 25 and the EuroQoL (EQ 5D 5L) questionnaire
3. Safety based on determined safety outcomes, adverse events, and serious adverse events
4. Number of treatments used (anti-VEGF injections, SML sessions) in the study eye from baseline to week 104
5. Number/proportion of people receiving “rescue” treatment in the study eye from baseline to week 104
6. Number of rescue treatments received in the study eye from baseline to week 104
7. Number/proportion of people discontinuing treatment (with reasons)
8. Number/proportion of people losing (with reasons)  $\geq 5$ ,  $\geq 10$  and  $\geq 15$  ETDRS letters of best-corrected visual acuity (from baseline to week 104) in the study eye
9. Number/proportion of people gaining  $\geq 5$ ,  $\geq 10$  and  $\geq 15$  ETDRS letters (from baseline to week 104) in the study eye
10. Number/proportion of people with CRT  $\leq 300\mu\text{m}$  in the study eye in the central 1 mm if the retina as determined using SD-OCT
11. Number/proportion of people with no DMO, as determined by the ophthalmologists evaluating the patient
12. Health and social care service use and non-healthcare costs as determined using a Health Service Use Questionnaire and Patient Cost Questionnaire
13. Participant experience and acceptability as determined by focus group discussions, the Acceptability Questionnaire ( Theoretical Framework of Acceptability (TFA) ) distributed at week 104, and also by the use of Visual Analogue Score questionnaires that will be distributed 60 minutes prior to treatment, immediately after treatment and 24 hours after treatment at all instances in which treatment is given

## **Completion date**

30/09/2028

## **Eligibility**

### **Key inclusion criteria**

1. Adults (>18 years)
2. Diabetes type 1 or type 2
3. Presented with severe centre-involving (CI)-DMO (CRT  $\geq 400\mu\text{m}$ )
4. Within the first year of initiating anti-VEGF therapy but who still have DMO and their CRT is below  $400\mu\text{m}$  (and it remains, at the time of randomisation) following anti-VEGF therapy in either one eye or both eyes

### **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. Causes of macular oedema other than DMO
2. DMO with CRT  $\geq 400$   $\mu\text{m}$
3. Receipt of anti-VEGFs before their presentation with severe DMO (previous macular laser treatment for DMO is allowed)
4. Use of unlicensed anti-VEGFs (e.g. bevacizumab)
5. Inability, for any reason, to attend study visits
6. Active proliferative diabetic retinopathy (PDR) (treated and inactive PDR is allowed)
7. Use of pioglitazone which cannot be stopped for the duration of the trial
8. Cataract surgery or laser pan-retinal photocoagulation (PRP) within the previous 6 weeks
9. Currently enrolled in a CTIMP (Clinical Trial of an Investigational Medical Product)
10. Declined consent for participation

**Date of first enrolment**

19/05/2025

**Date of final enrolment**

30/04/2026

**Locations****Countries of recruitment**

United Kingdom

England

Northern Ireland

Scotland

Wales

**Study participating centre**

**The Hillingdon Hospital**

Field Heath Road

Uxbridge

England  
UB8 3NN

**Study participating centre**  
**Frimley Park Hospital**  
Portsmouth Road  
Frimley  
Camberley  
England  
GU16 7UJ

**Study participating centre**  
**Kings College Hospital**  
Mapother House  
De Crespigny Park  
Denmark Hill  
London  
England  
SE5 8AB

**Study participating centre**  
**James Cook University Hospital**  
Marton Road  
Middlesbrough  
England  
TS4 3BW

**Study participating centre**  
**Chelsea and Westminster Hospital**  
Chelsea & Westminster Hospital  
369 Fulham Road  
London  
England  
SW10 9NH

**Study participating centre**  
**Central Middlesex Hospital**  
Acton Lane  
London  
England  
NW10 7NS

**Study participating centre**  
**Moorfields Eye Hospital**  
162 City Road  
London  
England  
EC1V 2PD

**Study participating centre**  
**Sunderland Eye Hospital**  
Queen Alexandra Rd  
Sunderland  
England  
SR2 9HP

**Study participating centre**  
**Royal Gwent Hospital**  
Cardiff Road  
Newport  
Wales  
NP20 2UB

**Study participating centre**  
**Bristol Eye Hospital**  
Lower Maudlin Street  
Bristol  
England  
BS1 2LX

**Study participating centre**  
**University Hospital Southampton**  
Southampton University Hospital  
Tremona Road  
Southampton  
England  
SO16 6YD

**Study participating centre**

**Gloucestershire Royal Hospital**  
Great Western Road  
Gloucester  
England  
GL1 3NN

**Study participating centre**  
**Royal Liverpool University Hospital**  
Royal Liverpool University Hospital  
Prescot Street  
Liverpool  
England  
L7 8XP

**Study participating centre**  
**Hull Royal Infirmary**  
Anlaby Road  
Hull  
England  
HU3 2JZ

**Study participating centre**  
**Queens Medical Centre**  
Nottingham University Hospital  
Derby Road  
Nottingham  
England  
NG7 2UH

**Study participating centre**  
**Royal Victoria Hospital**  
274 Grosvenor Road  
Belfast  
Northern Ireland  
BT12 6BA

**Study participating centre**  
**The Sussex Eye Hospital**  
Eastern Road

Brighton  
England  
BN2 5BF

**Study participating centre**

**Torbay Hospital**

Torbay Hospital  
Newton Road  
Torquay  
England  
TQ2 7AA

**Study participating centre**

**Birmingham Midland Eye Centre (bmec)**

City Hospital N H S Trust  
Dudley Road  
Birmingham  
England  
B18 7QH

**Study participating centre**

**Sandwell and West Birmingham Hospitals NHS Trust**

Midland Metropolitan University Hos  
Grove Lane  
Smethwick  
England  
B66 2QT

**Study participating centre**

**Singleton Hospital**

Sketty Lane  
Sketty  
Swansea  
Wales  
SA2 8QA

**Study participating centre**

**Swansea Bay University Local Health Board**

Tonna Hospital  
Tonna Uchaf  
Tonna

Neath  
Wales  
SA11 3LX

**Study participating centre**  
**Moorfields Eye Centre at Bedford Hospital (south)**  
Kempston Road  
Bedford  
England  
MK42 9DJ

**Study participating centre**  
**Whipps Cross Hospital**  
Leytonstone  
Whipps Cross Road  
London  
England  
E11 1NR

## **Sponsor information**

**Organisation**  
Belfast Health and Social Care Trust

**ROR**  
<https://ror.org/02tdmfk69>

## **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 and/or analysed during the current study during this study will be included in the subsequent results publication.

### IPD sharing plan summary

Published as a supplement to the results publication

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Protocol file</a>	version 2.0	09/09/2024	01/10/2024	No	No
<a href="#">Protocol file</a>	version 4.0	02/05/2025	30/12/2025	No	No
<a href="#">Study website</a>		11/11/2025	11/11/2025	No	Yes