# Extended follow-up of participants in the DROPLET randomised controlled trial

Submission date	Recruitment status  No longer recruiting	<ul><li>Prospectively registered</li></ul>		
13/03/2019		☐ Protocol		
Registration date	Overall study status	Statistical analysis plan		
18/04/2019	Completed	[X] Results		
Last Edited	Condition category	[] Individual participant data		
01/10/2021	Nutritional, Metabolic, Endocrine			

#### Plain English summary of protocol

Background and study aims

Around 1 in 4 people in the UK are seriously overweight (obese). Being obese makes it more likely that a person will develop diabetes, heart disease and some types of cancer. Extra body weight also puts a strain on muscles and joints, making it difficult and sometimes painful to move around. The aim of the DROPLET study, conducted in 2015-2017 was to investigate whether a GP referral of patients who are obese to a commercial low-energy total diet replacement (TDR) programme, could help these people to lose more weight than a weight loss programme provided by the practice nurse. 278 participants were recruited from GP practices across Oxfordshire CCG. The participants who were assigned to the TDR programme lost on average 10.7 kg after one year, compared with those assigned to the nurse advice who lost on average 3.1 kg, which resulted in an average difference between the groups of 7.2 kg in favour of the TDR group. These results clearly demonstrate that TDR is clinically effective at 1 year, and the health benefits are largely commensurate with the weight losses observed. However, there are concerns that a rapid weight may be followed with rapid regain, which could negate some of the health benefits. A better understanding of the longer-term effects on weight and health outcomes will help to determine whether the treatment is a cost-effective option in the longer term. The aim of this study is to contact the original participants in the DROPLET trial to invite them to

enrol in this study and to attend a new appointment about 3 years from their original randomisation visit to measure body weight and collect other information on their health and weight control efforts.

Who can participate?

Adults who participated in the original DROPLET study

What does the study involve?

Participants are seen by a nurse or a member of the research team to collect measures of weight, body fat, waist circumference and a fasting blood sample to assess change in blood markers, such as cholesterol and insulin. This is to assess participants' risk of developing heart disease or diabetes. Participants are also asked to complete questionnaires about weight control strategies. These measures collected at 3 years are compared with the same measures that were collected at the beginning of and during the main trial.

What are the possible benefits and risks of participating?

The risks of taking part are minimal, some people feel faint and have a small bruise where the blood sample has been collected.

Where is the study run from?
Oxfordshire Clinical Commissioning Group (UK)

When is the study starting and how long is it expected to run for? March 2019 to September 2019

Who is funding the study?

- 1. Cambridge Weight Plan Ltd (UK)
- 2. National Institutes of Health Research Collaboration for Leadership in Health Research and Care (UK)

Who is the main contact? Dr N Astbury nerys.astbury@phc.ox.ac.uk

# Contact information

#### Type(s)

Public

#### Contact name

Dr Nerys Astbury

#### **ORCID ID**

http://orcid.org/0000-0001-9301-7458

#### Contact details

University Of Oxford
The Radcliffe Observatory Quarter Woodstock Road
Nuffield Department of Primary Care Health Sciences
Oxford
United Kingdom
OX2 6GG
+44 (0)1856617871
nerys.astbury@phc.ox.ac.uk

### Additional identifiers

#### **EudraCT/CTIS** number

Nil known

**IRAS** number

#### ClinicalTrials.gov number

Nil known

#### Secondary identifying numbers

13987

# Study information

#### Scientific Title

Participants enrolled in the DROPLET randomised controlled trial which compared referral to a commercial total diet replacement (TDR) weight loss programme with usual care weight management on weight at 3 years

#### Study objectives

The aim of the Extended follow up of the Doctor Referral of Overweight People to a Low-Energy Treatment (DROPLET) trial (https://www.isrctn.com/ISRCTN75092026) is to determine the clinical effectiveness, of referral to a low-energy total diet replacement programme compared with usual weight management interventions for long term weight management.

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

South Central Oxford B Research Ethics Committee, Whitefriars, Level 3, Block B, Lewin's Mead, Bristol, BS1 2NTTel: +44 (0)207 1048058, Email: nrescommittee.southcentral-oxfordb@nhs.net, ref: 19/SC/0012

#### Study design

Observational follow-up study

#### Primary study design

Observational

#### Secondary study design

Cohort study

#### Study setting(s)

GP practice

#### Study type(s)

Treatment

#### Participant information sheet

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

Obesity

#### **Interventions**

This is an extended follow-up of the participants who were enrolled in the DROPLET randomised controlled trial (https://www.isrctn.com/ISRCTN75092026). Eligible participants have already taken part in the Doctor Referral of Overweight People to Low-Energy treatment (DROPLET) study whereby they were randomly assigned to either a TDR programme or usual care for weight loss, and were followed up at 1 year after randomisation. This study will invite those

participants to attend an extended follow-up at 3 years after randomisation. There will be no further randomisation or additional intervention; this is an observational follow-up only. The participants attend a new appointment approximately 3 years from their original randomisation visit to measure body weight and collect other information on their health and weight control efforts.

#### Intervention Type

Behavioural

#### Primary outcome measure

Weight change from baseline to 3 years

#### Secondary outcome measures

Proportion of participants in each group achieving 5% weight loss at 3 years Proportion of participants in each group achieving 10% weight loss at 3 years Change in fat mass between baseline and 3 years Change in LDL cholesterol between baseline and 3 years Change in HbA1c between baseline and 3 years Change in systolic and diastolic blood pressure between baseline and 3 years Change in QRISK2 between baseline and 3 years

#### Overall study start date

01/03/2019

#### Completion date

30/09/2019

# **Eligibility**

#### Key inclusion criteria

Participant in the original DROPLET randomised controlled trial and consented to re-contact

#### Participant type(s)

Other

#### Age group

Adult

#### Sex

Both

#### Target number of participants

272

#### Total final enrolment

179

#### Key exclusion criteria

Does not meet inclusion criteria

#### Date of first enrolment

01/03/2019

#### Date of final enrolment

30/09/2019

#### Locations

#### Countries of recruitment

England

**United Kingdom** 

# Study participating centre University of Oxford

The Radcliffe Observatory Quarter Woodstock Road Oxford United Kingdom OX2 6GG

# Sponsor information

#### Organisation

University of Oxford

#### Sponsor details

Clinical Trial and Research Governance
Joint Research Office
1st floor, Boundary Brook House
Churchill Drive
Headington
Oxford
England
United Kingdom
OX3 7GB
+44 (0)1865 289885
ctrg@admin.ox.ac.uk

#### Sponsor type

University/education

#### Website

https://researchsupport.admin.ox.ac.uk/ctrg

#### **ROR**

https://ror.org/052gg0110

# Funder(s)

#### Funder type

Industry

#### Funder Name

Cambridge Weight Plan UK Ltd

#### Funder Name

NIHR CLAHRC Oxford

# **Results and Publications**

#### Publication and dissemination plan

Planned publication in a high-impact peer reviewed journal sometime in 2020.

#### Intention to publish date

01/05/2020

#### Individual participant data (IPD) sharing plan

The current IPD data sharing plans for the current study are unknown and will be made available at a later date.

#### IPD sharing plan summary

Data sharing statement to be made available at a later date

#### **Study outputs**

Output type	<b>Details</b> results	Date created	Date added	Peer reviewed?	Patient-facing?
Results article		23/07/2021	01/10/2021	Yes	No
HRA research summary			28/06/2023	No	No