# Prevention of diabetes using structured education and continuous support programme in those with pre-diabetes in a multi-ethnic population

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
06/06/2008		[X] Protocol		
Registration date 11/07/2008	Overall study status Completed	Statistical analysis plan		
		[X] Results		
Last Edited	Condition category	[] Individual participant data		
23/03/2018	Signs and Symptoms			

# Plain English summary of protocol

Not provided at time of registration

# Contact information

# Type(s)

Scientific

### Contact name

**Prof Melanie Davies** 

## Contact details

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# Additional identifiers

**EudraCT/CTIS** number

IRAS number

ClinicalTrials.gov number

# Secondary identifying numbers

**UHLPRE01** 

# Study information

# Scientific Title

Prevention of diabetes using structured education and continuous support programme in those with pre-diabetes in a multi-ethnic population

# Acronym

**PREVENTION** 

# Study objectives

Structured education and continuous support programme reduce the incidence of diabetes in a multi-ethnic population with pre-diabetes.

# Ethics approval required

Old ethics approval format

# Ethics approval(s)

Not provided at time of registration

# Study design

Parallel controlled cluster randomised trial

# Primary study design

Interventional

# Secondary study design

Cluster randomised trial

# Study setting(s)

GP practice

# Study type(s)

Prevention

# Participant information sheet

Not available in web format, please use the contact details below to request a patient information sheet

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

Pre-diabetes

# Interventions

Intervention arm:

Six hours of education within one month of diagnosis, either as one full day, two half days, or 4 x 2 hours (delivered with interpreters for participants whose first language is not english); follow-

up will be for three years with clinical assessments at 6, 12, 24, 23 months. Continuous support will be regular telephone contact, website and additional written material.

### Control arm:

Standard care as per national guidelines currently general lifestyle advice and written material follow-up will be for three years with clinical assessments at 6, 12, 24, 23 months.

# Intervention Type

Behavioural

# Primary outcome measure

Reduction in the incidence of diabetes at three years

# Secondary outcome measures

- 1. Reduction in cardiovascular disease (CVD) risk (assessed using the current Framingham risk equation, standard for CVD risk assessment for General Practice in the UK), measured at baseline and 6, 12, 24, 23 months
- 2. Reduction in HbA1c, fasting and two-hour glucose, measured at baseline and 6, 12, 24, 23 months

# Overall study start date

01/09/2008

# Completion date

31/12/2012

# **Eligibility**

# Key inclusion criteria

- 1. Aged 40 70 years (if White European)
- 2. Aged 25 70 years (if South Asian)
- 3. Able to attend group education sessions

# Participant type(s)

All

# Age group

Adult

## Sex

Both

# Target number of participants

816

# Key exclusion criteria

- 1. Unable to give informed consent
- 2. Diagnosis of diabetes

## Date of first enrolment

# Date of final enrolment 31/12/2012

# Locations

# Countries of recruitment

England

**United Kingdom** 

Study participating centre Leicester Diabetes Centre (Broadleaf)

Leicester United Kingdom LE5 4PW

# Sponsor information

# Organisation

University Hospitals of Leicester NHS Trust (UK)

# Sponsor details

Trust Headquarters Gwendolyn House Gwendolyn Road Leicester England United Kingdom LE5 4QF

# Sponsor type

Hospital/treatment centre

# Website

http://www.uhl-tr.nhs.uk/

## **ROR**

https://ror.org/02fha3693

# Funder(s)

# Funder type

Government

# Funder Name

National Institute of Health Research (NIHR) (UK) - Programme Grant for Applied Research (PGFAR) (ref: RG-PG-0606-1272)

# **Results and Publications**

# Publication and dissemination plan

Not provided at time of registration

Intention to publish date

Individual participant data (IPD) sharing plan

# IPD sharing plan summary

Not provided at time of registration

# **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol article	protocol	20/05/2012		Yes	No
Results article	screening results	23/12/2014		Yes	No
Results article	baseline results	23/07/2015		Yes	No