# Diabetes 6 (D-6) trial: psychologically enhanced diabetes care, delivered by nurses

<b>Submission date</b> 19/05/2010	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered
		☐ Protocol
Registration date 19/05/2010	Overall study status Completed	Statistical analysis plan
		[X] Results
Last Edited	Condition category	[] Individual participant data
14/06/2017	Nutritional, Metabolic, Endocrine	

#### Plain English summary of protocol

Not provided at time of registration

# Contact information

# Type(s)

Scientific

#### Contact name

Dr Kirsty Winkley

#### Contact details

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# Additional identifiers

**EudraCT/CTIS** number

IRAS number

ClinicalTrials.gov number

**Secondary identifying numbers** 4299

# Study information

#### Scientific Title

A randomised controlled trial of 'Psychologically Enhanced Diabetes Care, delivered by nurses' (D-6) for people with Type 2 diabetes and sub-optimal glycaemic control

#### Acronym

D-6

#### Study objectives

Some people with type 2 diabetes (T2DM) have problems reaching ideal blood glucose levels that will prevent a worsening of their condition. They may need to make significant lifestyle changes to do this which can be very difficult. Often it may be a question of being unsure whether they can make the changes not that they don't know what changes to make. Talking therapies such as motivational interviewing and cognitive behavioural therapy can help people talk through their difficulties within a supportive relationship and help foster their own self confidence in their ability to make changes. These therapies have increasingly been used in health settings although usually delivered by psychologists but the skills involved can be taught to non-psychologists such as practice nurses.

The aim of this study is to train practice nurses (for intervention GP practices) in talking therapy for diabetes and compare their patients with those from practices where nurses have not had the training. We would invite approximately 24 GP practices from the London boroughs of Lambeth, Southwark and Lewisham to participate and these would be randomised and nurses in the intervention practices would receive training. Nurses working in the non-intervention (control) GP practices would receive additional training in diabetes education. If this study shows that training nurses in talking therapy helps their patients to get better control of their diabetes this training could be offered to other practice nurses working in the NHS.

### Ethics approval required

Old ethics approval format

# Ethics approval(s)

- 1. South East Ethics Committee 3, 30/11/2009, ref: 09/H0808/97
- 2. Research & Development Centre for South East London NHS, 05/05/2010, ref: RDLSL 534/CSP 20423

# Study design

Multicentre randomised interventional treatment trial

# Primary study design

Interventional

#### Secondary study design

Randomised controlled trial

# Study setting(s)

GP practice

#### Study type(s)

Treatment

#### Participant information sheet

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

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

Topic: Diabetes Research Network; Subtopic: Type 2; Disease: Hypertension, Dyslipidaemia, Diabetic Control, Obesity, Prevention/screening

#### **Interventions**

Baseline blood tests are taken in both the treatment and control groups. Patients are randomised to:

- 1. D-6 treatment group: face to face psychological diabetes care sessions. Maintenance psychological diabetes sessions, specified by patient, may be telephone/e-mail/face to face.
- 2. Control group: Face to face diabetes education sessions. Maintenance education sessions specified by patient, may be telephone/e-mail/face to face.

Follow-up blood tests are then taken in both the treatment and control groups.

Total duration of intervention: 12 months Total duration of follow-up: 6 months

#### Intervention Type

Other

#### **Phase**

Phase II

#### Primary outcome measure

Change in HbA1c (glycaemic control) at 18 months

#### Secondary outcome measures

- 1. Total cholesterol, taken at baseline and at end of study
- 2. Blood pressure, taken at baseline and at end of study
- 3. BMI, taken at baseline and at end of study
- 4. Quality of life, measured before and after intervention
- 5. Depressive symptoms, measured before and after intervention
- 6. Economic evaluation, measured before and after intervention

#### Overall study start date

15/10/2007

#### Completion date

30/11/2016

# **Eligibility**

#### Key inclusion criteria

- 1. People with Type 2 diabetes for more than 2 years
- 2. Resident in south London boroughs of Lambeth Southwark and Lewisham

- 3. Aged 18 79 years, either sex
- 4. Persistent suboptimal glycaemic control defined as HbA1c above or equal to 8% on occasions in the past 12 months despite 2 appointments with a diabetes health professional

#### Participant type(s)

Patient

#### Age group

Adult

#### Lower age limit

18 Years

#### Sex

Both

#### Target number of participants

Planned sample size: 432

#### Key exclusion criteria

- 1. Severe mental disorders
- 2. Terminal illnesses and severe end stage diabetes complications
- 3. Morbid obesity with a body mass index (BMI) greater than 50 kg/m^2 as there is usually comorbid psychiatric morbidity
- 4. No telephone or mobile phone access as telephone contact is part of the intervention

#### Date of first enrolment

27/09/2010

#### Date of final enrolment

11/10/2011

# Locations

## Countries of recruitment

England

United Kingdom

# Study participating centre Department of Psychological Medicine

London United Kingdom SE5 9RJ

# Sponsor information

#### Organisation

King's College Hospital NHS Foundation Trust (UK)

#### Sponsor details

Department of Research & Development 34 Love Walk London England United Kingdom SE5 8AD

#### Sponsor type

Hospital/treatment centre

#### Website

http://www.kch.nhs.uk/

#### **ROR**

https://ror.org/01n0k5m85

# Funder(s)

# Funder type

Government

#### **Funder Name**

National Institute for Health Research (NIHR) (UK) - Programme Grants for Applied Research (ref: RP-PG-0606-1142)

# **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?

Results article results 01/10/2016 Yes

No