Project CYGNUS: A feasibility study of gathering health information in people with memory problems

Submission date	Recruitment status	Prospectively registered		
27/03/2017	No longer recruiting	☐ Protocol		
Registration date	Overall study status	Statistical analysis plan		
19/04/2017	Completed	Results		
Last Edited	Condition category	Individual participant data		
22/05/2019	Mental and Behavioural Disorders	Record updated in last year		

Plain English summary of protocol

Background and study aims

Alzheimer's disease (AD) is a common cause of dementia. People with dementia have difficulties with mental processes such as memory, language, reasoning and identifying people and objects, which become progressively worse over time as they age. Recently, there have been failures in developing medications to try and cure AD. This means new efforts are needed to support those suffering without the hope of a cure. New projects are being undertaken that are focused on aiding studies of new AD treatments. While these research goals are important, they are part of a strategy that will not impact patients for over 15 years. The aim of this study is to explore how the information collected at Memory Assessment Services when patients are referred and diagnosed with AD and/or dementia and additional information collected from patients and their study partners over a one year follow-up period could be used to improve patient treatment and care.

Who can participate?

Adults over the age of 18 who are referred to a Memory Assessment Service and their partner, family member or close friend.

What does the study involve?

Participants are visited in their own homes (or in a clinic if they prefer) where they are asked a series of questionnaires about living with dementia. Participants are either patients who have been referred to memory assessment services or study partners who are those who have consented to support the patient through the study and are happy to answer a set of questionnaires aimed at carers of people living with dementia. This takes place every three months for one year. Participants are also asked if they would like to use a wearable device that collects data about activity and sleep over two or 12 weeks (depending on the device they use).

What are the possible benefits and risks of participating? There are no notable benefits or risks with participating.

Where is the study run from?

This study is being run from the University of Manchester (UK) and takes place in NHS Mental Health Trusts across England (UK).

When is the study starting and how long is it expected to run for? April 2016 to March 2018

Who is funding the study? Innovate UK (UK)

Who is the main contact?

1. Ms Francine Jury (Public)
francine.jury@manchester.ac.uk

2. Professor Derek Hill (Scientific)

Study website

ecygnus.com

Contact information

Type(s)

Public

Contact name

Ms Francine Jury

Contact details

Room 3.306
Jean McFarlane Building
University of Manchester
Oxford Road
Manchester
United Kingdom
M13 9PY
+44 161 306 7491
francine.jury@manchester.ac.uk

Type(s)

Scientific

Contact name

Prof Derek Hill

Contact details

Ixico PLC 4th Floor Griffin Court 15 Long Lane London United Kingdom EC1A 9PN

Additional identifiers

EudraCT/CTIS number

IRAS number

ClinicalTrials.gov number

Secondary identifying numbers 30502

Study information

Scientific Title

Project CYGNUS: A feasibility study of gathering health information in people with memory problems

Study objectives

The aim of this study is to explore how the information collected at Memory Assessment Services at the point of referral and subsequent diagnosis and additional information collected from the cognitively impaired participants and their study partners over a one year follow-up period could be used to improve patient treatment and care.

Ethics approval required

Old ethics approval format

Ethics approval(s)

London – Central Research Ethics Committee, 23/04/2016, ref: 16/LO/0354

Study design

Observational; Design type: Cohort study

Primary study design

Observational

Secondary study design

Cohort study

Study setting(s)

Hospital

Study type(s)

Treatment

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

Specialty: Dementias and neurodegeneration, Primary sub-specialty: Dementia; UKCRC code/ Disease: Neurological/ Other degenerative diseases of the nervous system

Interventions

Participants receive five home visits at three month intervals over 12 month periods. This involves being asked a series of questionnaires relating to outcomes for people living with dementia (the standard scales for dementia). The questions are answered by either the participant or by their study partners who have consented to support the participant through the study and are happy to answer a set of questionnaires aimed at carers of people living with dementia.

Participants are also asked if they would like to take part in a sub-study. This includes either using a wearable device to collect continuous data on activity and sleep over two or 12 weeks (depending on the device) or using mobile data collection which involves using a web/mobile app to collect self-reported data on a more regular basis from home (this is done weekly).

There are 2 wearable devices are:

The Withings device: This device is worn for 12 weeks and collects sleep and activity data. It is a commercial device which requires other technology to use such as a smartphone or tablet that can download an app to collect self-reported data on a more regular basis from home (this is done weekly).

The Axivity AX3 device: This device is worn for 2 weeks and does not required any other technology. It collects raw data over two weeks which is then downloaded.

The data collected is reviewed to examine the feasibility of collecting quality and actionable patient and carer reported outcomes.

Intervention Type

Other

Primary outcome measure

Information about Memory Assessment Services are measured using patient and carer reported outcomes (PRO) (Quality of Life, Activates of Daily Living, etc) at baseline, three, six, nine and twelve months.

Secondary outcome measures

Feasibility to collect quality and actionable patient and carer reported outcomes is measured using web-based and mobile technologies continuously or at weekly increments.

Overall study start date

14/04/2016

Completion date

31/03/2018

Eligibility

Key inclusion criteria

Cognitively Impaired Participants:

- 1. Aged 18 years or older
- 2. Referred to a Memory Assessment Service with suspected Dementia by GP

- 3. Able to understand written and spoken English with the assistance of a carer/ study partner
- 4. Have capacity to consent

Study Partners:

- 1. Aged 18 years or older
- 2. Partner, Family Member or Close Friend is a participant in CYGNUS in the cognitively impaired participant group
- 3. English speaking/able to understand study documents

Participant type(s)

Patient

Age group

Adult

Lower age limit

18 Years

Sex

Both

Target number of participants

Planned Sample Size: 330; UK Sample Size: 330

Key exclusion criteria

Cognitively Impaired Participants:

- 1. Aged younger than 18 years
- 2. Currently resident in a care or nursing home
- 3. Do not have capacity to consent

Study Partners:

- 1. Aged younger than 18 years
- 2. Do not have capacity to consent
- 3. Previously referred to memory assessment service themselves

Date of first enrolment

01/04/2016

Date of final enrolment

31/03/2017

Locations

Countries of recruitment

England

United Kingdom

Study participating centre

University of Manchester

Room 3.306
Jean McFarlane Building
Oxford Road
Manchester
United Kingdom
M13 9PY

Sponsor information

Organisation

Ixico PLC

Sponsor details

4th Floor Griffin Court 15 Long Lane London United Kingdom EC1A 9PN

Sponsor type

Industry

Website

http://www.ixico.com

ROR

https://ror.org/00paezp73

Funder(s)

Funder type

Government

Funder Name

Innovate UK

Alternative Name(s)

innovateuk

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Results and Publications

Publication and dissemination plan

Planned publication in a high impact journal.

Intention to publish date

31/05/2019

Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are not expected to be made available due to the dataset is collected on a proprietary data platform at Ixico PLC and used for platform development. Paper based data collection could be made available on request from June 2018 but there would be a cost for retrieval for data archives.

IPD sharing plan summary

Not expected to be made available

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
HRA research summary			28/06/2023	No	No