

# Social care for people living with young onset dementia: the DYNAMIC study

<b>Submission date</b> 06/12/2023	<b>Recruitment status</b> No longer recruiting	<input type="checkbox"/> Prospectively registered <input checked="" type="checkbox"/> Protocol
<b>Registration date</b> 12/12/2023	<b>Overall study status</b> Completed	<input type="checkbox"/> Statistical analysis plan <input checked="" type="checkbox"/> Results
<b>Last Edited</b> 01/12/2025	<b>Condition category</b> Mental and Behavioural Disorders	<input type="checkbox"/> Individual participant data

## Plain English summary of protocol

### Background and study aims

The aim of this study is to identify, prioritise and produce recommendations and resources to address improvements in social care for people with young onset dementia (dementia that starts under the age of 65 years) and their families.

Social care for people with young onset dementia is seldom well addressed. Social care needs are very different from those of older people and young onset dementia also impacts families, including children/young people. Little research has been carried out in this area, with only six UK studies located in a 2018 review. The largest UK study of support for people with young onset dementia to date, the Angela project, found that social care is central to quality of life in young onset dementia. The researchers want to build on this by focusing on social care practice and how this can be improved.

### Who can participate?

1. People living with or family members/friends/neighbours supporting someone with young onset dementia in England will be eligible to take part in a face-to-face or online interview. The researchers will select from those who express an interest to make sure we hear diverse stories from a wide range of people.
2. Staff with a role in social care will be able to take part in the online survey from anywhere in England.
3. Up to 20 diverse stakeholders will be invited to take part in setting priorities and in the development of the resources and recommendations.

### What does the study involve?

People living with Young Onset Dementia or supporting someone with young onset dementia will take part in a one-to-one, 30-60 minute interview with a researcher about their situation and experiences of social care and to find out about their social care needs and support they would prefer.

Staff involved in social care for people with Young Onset Dementia will be invited to complete a 10-minute online survey to find out about their awareness, knowledge and practice.

Stakeholders involved in setting priorities and in producing recommendations and resources will be asked to attend a 90-minute online workshop and give a further 90 minutes to assist with feedback on materials. The stakeholders will be asked to agree on priority improvements.

Finally, the researchers will produce recommendations and resources that address these priority issues. These will be informed by talking with 10-12 professionals who work in existing pockets of excellence and by searching online for evidence about good practice in the relevant areas.

What are the possible benefits and risks of taking part?

People with dementia and carers may find it hard to make time for interviews. Staff are likely to be pressured at work and have little time for research. There is some risk of distress to people with young onset dementia and main supporters from discussing their experiences or situation. There is some risk that those taking part may disclose information which the researcher feels poses a serious risk to the participant or others. In this event, the researcher will discuss this with the research team and the team will contact the Local Authority Adult Safeguarding Team, if appropriate.

People with young onset dementia and supporters often report that it is beneficial for them to talk out loud about their experiences to someone who is attentively listening and interested. In addition, those taking part may gain satisfaction from contributing to a project that is about producing recommendations and resources to improve future social care for people with YOD and their families.

Where is the study run from?

University of Bradford (UK)

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

DYNAMIC started on 17/4/2023 and will run until 17/10/2025.

Who is funding the study?

1. National Institute for Health and Care Research (NIHR) (UK)
2. University of York (UK)
3. Dementia UK (UK)

Who is the main contact?

1. Prof. Jan Oyeboode, [j.oyebode@bradford.ac.uk](mailto:j.oyebode@bradford.ac.uk)
2. Dr Catherine Quinn, [c.quinn1@bradford.ac.uk](mailto:c.quinn1@bradford.ac.uk)

## Contact information

### Type(s)

Scientific, Principal investigator

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### **Type(s)**

Public

### **Contact name**

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## **Additional identifiers**

### **Clinical Trials Information System (CTIS)**

Nil known

### **Integrated Research Application System (IRAS)**

329325

### **ClinicalTrials.gov (NCT)**

Nil known

### **Protocol serial number**

IRAS 329325

## **Study information**

### **Scientific Title**

Social care planning and provision for people with young onset dementia and their families: current practice and resources for improvement

### **Acronym**

DYNAMIC

### **Study objectives**

Research has established the high level and breadth of social care needs in young onset dementia (YOD), the lack of social care for YOD and the deleterious consequences of living with unmet social care needs for the person, main supporter and children/young people. Consultation work with a range of stakeholders has indicated that no-one seems to have a clear picture of

current social care for YOD. This project aims to address this gap by exploring the social care experiences of people living with YOD and their families and finding out about current practice in social care provision. Based on these findings, a co-production approach with key stakeholders will be used to establish priorities for improvement, and produce 2-3 evidence-based recommendations and resources for improvement.

### **Ethics approval required**

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### **Ethics approval(s)**

1. approved 20/09/2023, Social Care REC Health Research Authority (2nd Floor, 2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 (0)207 104 8070; socialcare.rec@hra.nhs.uk), ref: 23/IEC08/0034

2. approved 01/08/2023, University of Bradford Research Ethics Panel (Research & Innovation Services (RAIS), F.24 Richmond Building, Richmond Road, Bradford, BD7 1DP, United Kingdom; +44 (0)1274 236554; nhs-ethics@bradford.ac.uk), ref: IRAS208

The researchers have received approval from the University of Bradford and the HRA Social Care REC for work packages 1 & 2 of the study. If required, they plan to submit a further application for ethical approval of work package 3 in a time-appropriate manner.

### **Study design**

Mixed methods single-centre observational study

### **Primary study design**

Observational

### **Study type(s)**

Quality of life, Other

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

Social care for people living with young onset dementia and their families

### **Interventions**

The DYNAMIC study involves three work packages. The first two will run in parallel over 15-16 months and be followed by the third over the subsequent 15 months.

In work package 1, the project will gather accounts, using in-depth qualitative interviews, from 25 people with young onset dementia and/or their family/friend carers to find out about social care needs, experiences and support they would prefer. Data will be subject to inductive thematic analysis.

In work package 2, a national 10-minute survey of staff with a role in social care will be conducted, to find out about their awareness, knowledge and practice of social care for people with young onset dementia and their families. This will be subject to descriptive data analysis and content analysis.

In work package 3, the findings to date will be presented to 'stakeholders' via a workshop to agree on priority improvements. A search for good practices in these priority areas will then be completed by looking at published evidence and interviewing professionals. Work will then be

undertaken using co-production to produce recommendations and/or resources to improve social care in 2-3 of the priority areas.

**Intervention Type**

Other

**Primary outcome(s)**

Experiences of social care for young onset dementia and areas for improvement established through thematic analysis and descriptive data measured by interview at a single time point

**Key secondary outcome(s)**

There are no secondary outcome measures

**Completion date**

17/10/2025

**Eligibility****Key inclusion criteria**

Work package 1 participants will be people with YOD and main supporters:

1. Living in England.
2. Diagnosed with YOD or a relative, friend or neighbour of someone diagnosed with YOD
3. Living alone or with others

Work package 2 participants will be staff with a role in or awareness of adult/older adult social care planning, provision, management or commissioning, including but not limited to:

1. Local authority social workers
2. Local authority community OTs
3. Local authority social services commissioners, managers and coordinators
4. Social prescribers from third sector and primary care organisations
5. Staff from third-sector organisations that provide community and/or residential services, e.g., dementia navigators or link workers, Young Onset Dementia Advisors, support group facilitators
6. Staff from private providers e.g. home carers, personal assistants, care home staff and managers

**Participant type(s)**

Carer, Patient, Other

**Healthy volunteers allowed**

No

**Age group**

Mixed

**Lower age limit**

18 years

**Upper age limit**

100 years

**Sex**

All

**Total final enrolment**

185

**Key exclusion criteria**

People with young onset dementia will be excluded if:

1. They are diagnosed with dementia caused by HIV, traumatic brain injury, Down's syndrome, Huntington's chorea or alcohol-related dementia
2. They are unable to communicate or lack capacity to consent to take part

**Date of first enrolment**

03/11/2023

**Date of final enrolment**

19/06/2025

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

United Kingdom

England

**Study participating centre****University of Bradford**

Centre for Applied Dementia Studies

Faculty of Health Studies

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**Sponsor information****Organisation**

University of Bradford

**ROR**

<https://ror.org/00vs8d940>

# Funder(s)

## Funder type

Government

## Funder Name

National Institute for Health and Care Research NIHR204266

## 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

## Funder Name

University of Bradford

## Alternative Name(s)

## Funding Body Type

Private sector organisation

## Funding Body Subtype

Universities (academic only)

## Location

United Kingdom

## Funder Name

Dementia UK

# Results and Publications

## Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study will be stored in a publicly available repository. Anonymised research data will be archived after the end of the

study using facilities provided by the UK Data Archive (<https://www.data-archive.ac.uk/>). Participants will be asked to consent for their data to be archived.

## IPD sharing plan summary

Stored in publicly available repository

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<a href="#">Results article</a>		20/09/2025	01/12/2025	Yes	No
<a href="#">Protocol article</a>		05/02/2024	06/02/2024	Yes	No
<a href="#">Other files</a>		06/10/2025	24/11/2025	No	No
<a href="#">Plain English results</a>	version 1	16/09/2025	20/10/2025	No	Yes
<a href="#">Plain English results</a>			24/11/2025	No	Yes
<a href="#">Study website</a>	Study website	11/11/2025	11/11/2025	No	Yes