

The HOME Study: A study comparing two different ways of looking after older people in hospitals to see if we can improve how quickly they go home

Submission date 03/01/2018	Recruitment status No longer recruiting	☒ Prospectively registered ☒ Protocol
Registration date 12/01/2018	Overall study status Completed	☒ Statistical analysis plan ☒ Results
Last Edited 18/12/2025	Condition category Other	☐ Individual participant data

Plain English summary of protocol

Background and study aims

NHS general hospitals have more than two million unplanned admissions of people aged 65 and older every year. These patients typically spend more time in hospital than those aged under 65. Long hospital stays are bad for older patients: they can get new illnesses like infections and lose their independence. They are also bad for the NHS which has a shortage of hospital beds.

Psychological problems, like dementia, confusion, depression and anxiety, are common in older patients and these are an important cause of long hospital stays. These problems are often not identified in busy hospital wards where the focus is on patients' physical illnesses. The aim of this study is to find out whether adding a new approach (sometimes called Proactive Liaison Psychiatry) to the identification and management of psychological problems reduces the time that older people spend in acute general hospital wards.

Who can participate?

Patients aged 65 and older who have been admitted to acute wards in hospitals in Oxfordshire, Cambridgeshire and Devon

What does the study involve?

Participants are randomly allocated to receive either usual care, or usual care plus the new approach (which involves seeing a doctor, nurse or occupational therapist who specialises in psychological problems in the medically ill). The study looks at whether the new approach reduces the time that patients spend in the hospital and whether it improves their quality of life and independence. Patients, carers and healthcare professionals are also interviewed to learn about their experiences with the new approach.

What are the possible benefits and risks of participating?

Taking part involves participants talking with the research team and allowing them access to their information. They may see an additional doctor or nurse who specialises in psychological problems. However, it is not known whether this will help them get home more quickly; that is

why this study is happening. The aim of the study is to find out how best to care for patients in the future.

Where is the study run from?

1. John Radcliffe Hospital (UK)
2. Horton Hospital (UK)
3. Royal Devon and Exeter Hospital (UK)
4. Addenbrooke's Hospital (UK)

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

April 2017 to October 2022

Who is funding the study?

National Institute for Health Research (NIHR) (UK)

Who is the main contact?

Dr Jane Walker, jane.walker@psych.ox.ac.uk (UK)

Contact information

Type(s)

Scientific

Contact name

Dr Jane Walker

ORCID ID

<https://orcid.org/0000-0003-1938-1141>

Contact details

Psychological Medicine Research Group
University of Oxford Department of Psychiatry
Oxford
United Kingdom
OX3 7JX

Additional identifiers

Central Portfolio Management System (CPMS)

36034

Study information

Scientific Title

The HOME Study: a study comparing two different ways of looking after older people in hospitals to see if we can improve how quickly they go home

Acronym

HOME

Study objectives

NHS general hospitals have more than two million unplanned admissions of people aged 65 and older every year. These patients typically spend more time in hospital than those aged under 65. Long hospital stays are bad for older patients: they can get new illnesses like infections and lose their independence. They are also bad for the NHS which has a shortage of hospital beds. Psychological problems, like dementia, confusion, depression and anxiety, are common in older patients and these are an important cause of long hospital stays. These problems are often not identified in busy hospital wards where the focus is on patients' physical illnesses.

In this study, the trialists will research whether adding a new approach (sometimes called Proactive Liaison Psychiatry) to the identification and management of psychological problems reduces the time that older people spend in acute general hospital wards.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved 02/11/2017, Oxford C REC (Health Research Authority (Bristol), Ground Floor, Temple Quay House, 2 The Square, BS1 6PN, Bristol, UK; +44 (0)207 1048144, (0)207 1048241, (0)207 1048289; oxfordc.rec@hra.nhs.uk), ref: 17/SC/0497

Primary study design

Interventional

Study design

Randomized complex intervention with qualitative component study

Study type(s)

Treatment

Health condition(s) or problem(s) studied

Older people admitted to hospital

Interventions

The trialists will recruit approximately 3,588 patients aged 65 and older, who have been admitted to acute wards in hospitals in Oxfordshire, Cambridgeshire and Devon. Half of the study participants will get usual NHS care. This means that they will be looked after by doctors, nurses and other staff in the usual way during their hospital stay. The other half will get usual NHS care and they will also see a doctor or nurse who specialises in psychological problems – they will talk to them to find out if they have problems such as worries or confusion that might delay their discharge from hospital and will make a plan to help them with these. The trialists will study whether the new approach reduces the time that patients spend in hospital and whether it improves their quality of life and independence. They will also interview patients, carers and healthcare professionals to learn about their experiences of the new approach.

Intervention Type

Other

Primary outcome(s)

Number of days spent as an inpatient in a general hospital in the month (30 days) post-randomisation; Timepoint(s): 30 days

Key secondary outcome(s)

Current secondary outcome measures as of 17/10/2022:

1. Cognitive function, measured using MOCA-T at 1 and 3 months
2. Independent functioning, measured using Barthel Index of Activities of Daily Living at 1 and 3 months
3. Health-related quality of life, measured using EQ-5D-5L at 1 and 3 months
4. Symptoms of anxiety and depression, measured using PHQ-4 at 1 and 3 months
5. Overall quality of life, measured using study-specific item at 1 and 3 months
6. Patient's experience of hospital stay, measured using study-specific item at 1 month
7. Patient's view on the length of their hospital stay, measured using study-specific item at 1 month
8. Discharge destination
9. Secondary healthcare use (including total length of index admission, length of index admission truncated at 30 days, number of readmissions, number of days in hospital) in the year post-randomisation
10. Death in the year post-randomisation

Previous secondary outcome measures as of 15/02/2018:

1. Cognitive function, measured using MOCA-T at 1 and 3 months
2. Independent functioning, measured using Barthel Index of Activities of Daily Living at 1 and 3 months
3. Health-related quality of life, measured using EQ-5D-5L at 1 and 3 months
4. Symptoms of anxiety and depression, measured using PHQ-4 at 1 and 3 months
5. Overall quality of life, measured using study-specific item at 1 and 3 months
6. Patient's experience of hospital stay, measured using study-specific item at 1 month
7. Patient's view on the length of their hospital stay, measured using study-specific item at 1 month
8. Discharge destination
9. Secondary healthcare use (including total length of index admission, number of readmissions, number of days in hospital) in the year post-randomisation
10. Death in the year post-randomisation

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7. Patient's view on the length of their hospital stay, measured using study-specific item at 1 and 3 months
8. Discharge destination
9. Secondary healthcare use (including total length of index admission, number of readmissions, number of days in hospital) in the year post-randomisation
10. Death in the year post-randomisation

Completion date

31/10/2022

Eligibility

Key inclusion criteria

To be included in the study patients must:

1. Be an inpatient in an acute ward
2. Have been admitted non-electively
3. Be aged 65 years or older
4. Be expected (by their clinical team) to remain an inpatient for at least 2 days from the time of baseline assessment
5. Be able to give informed consent or if unable to give consent, a consultee advises that study participation is appropriate

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Senior

Lower age limit

65 Years

Upper age limit

100 Years

Sex

All

Total final enrolment

2744

Key exclusion criteria

Patients will be excluded if:

1. They are moribund (defined as when the clinicians caring for a patient estimate that they are likely to die before discharge from hospital)
2. Their participation in the study is judged to be clinically or practically inappropriate (e.g. patient visiting from overseas)
3. They have already been enrolled in the study
4. They have already been referred to the usual liaison psychiatry team.
5. They have been a general hospital inpatient continuously for 1 week at the time of the baseline assessment
6. They do not read or speak English

Date of first enrolment

12/02/2018

Date of final enrolment

05/03/2020

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

John Radcliffe Hospital

Headley Way

Headington

Oxford

England

OX3 9DU

Study participating centre

Horton General Hospital

81a Oxford Road

Banbury

England

OX16 9AL

Study participating centre

Royal Devon and Exeter Hospital

Gladstone Road

Exeter

England

EX1 2ED

Study participating centre

Addenbrookes

Addenbrookes Hospital

Hills Road

Cambridge

England

CB2 0QQ

Sponsor information

Organisation
University of Oxford

ROR
<https://ror.org/052gg0110>

Funder(s)

Funder type
Government

Funder Name
NIHR Evaluation, Trials and Studies Co-ordinating Centre (NETSCC); Grant Codes: 15/11/16

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary
Not expected to be made available

Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Results article		10/08/2024	12/08/2024	Yes	No
Results article		01/12/2025	18/12/2025	Yes	No
Protocol article	protocol	07/08/2019	27/08/2019	Yes	No
HRA research summary			28/06/2023	No	No
Participant information sheet	version V2.0	17/10/2017	12/01/2018	No	Yes
Statistical Analysis Plan	Statistical and economic analysis plan	04/05/2020	06/05/2020	No	No
Study website		11/11/2025	11/11/2025	No	Yes