

Optimising the feasibility and acceptability of a multi-component, digital health intervention to improve outcomes for people with chronic obstructive pulmonary disease

Submission date 05/06/2019	Recruitment status No longer recruiting	☐ Prospectively registered ☒ Protocol
Registration date 05/07/2019	Overall study status Completed	☐ Statistical analysis plan ☒ Results
Last Edited 28/01/2026	Condition category Respiratory	☐ Individual participant data

Plain English summary of protocol

Background and study aims

Digital healthcare can address a number of health care needs arising from long-term conditions, including the requirement to support people in their own homes and to provide a greater focus on self-management. Substantial amounts of data are routinely captured during NHS usual care, including vital sign data (such as blood pressure and pulse rate) whilst patients are in hospital. Linking data routinely collected in hospital with data collected by monitoring devices at home has the potential to provide a better overview of health status, but the feasibility of doing this is currently unknown. Large numbers of admissions associated with an exacerbation of chronic obstructive pulmonary disease (COPD) are recorded each year with 1 in 4 patients re-admitted within 3 months of discharge. Linking data from these sources may help inform subsequent studies in developing more accurate predictive algorithms to identify deterioration and prevent readmission.

Who can participate?

Patients aged 40 years or older diagnosed with COPD who are a current or ex-smoker.

What does the study involve?

EDGE2 will recruit 200 patients diagnosed with chronic obstructive pulmonary disease (COPD). Patients will be approached if they visit hospital because of their COPD. A COPD exacerbation is a severe worsening of symptoms beyond usual day-to-day variation and can be a frightening experience. Participants will be given a tablet computer and two monitors to track oxygen saturation levels, heart rate and step count. Patients will access these devices whilst in hospital and for 6 months after discharge. EDGE2 will investigate the feasibility of accessing hospital data and see if it can be linked to self-monitored data. Linking this data together is important to develop predictive algorithms to identify and support responses to early changes in their health. EDGE2 will also monitor patients for 5 years after hospital discharge to identify hospital visits and deaths using data from NHS Digital.

What are the possible benefits and risks of participating?

Participants will be able to self-monitor their vital signs (including oxygen saturation), COPD symptoms, mood and physical activity on a regular basis over several months. They will also be able to access educational resources to support their COPD self-management, such as videos to demonstrate correct inhaler technique. In doing so participants may feel more anxious about their COPD by monitoring their condition more closely or may consider this a replacement to usual care. Incoming data will, however, be monitored by a member of the clinical research team twice-weekly and participants will also be able to contact the team themselves.

Where is the study run from?

Oxford University Hospitals NHS Foundation Trust, UK.

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

May 2019 to January 2021

Who is funding the study?

1. National Institute for Health Research Oxford Biomedical Research Centre
2. Engineering and Physical Sciences Research Council

Who is the main contact?

1. Prof. Andrew Farmer,
andrew.farmer@phc.ox.ac.uk
2. Miss Bethany Lawson,
bethany.lawson@phc.ox.ac.uk

Contact information

Type(s)

Scientific

Contact name

Prof Andrew Farmer

ORCID ID

<https://orcid.org/0000-0002-6170-4402>

Contact details

Department of Primary Care Health Sciences
Radcliffe Observatory Quarter
Woodstock Road
Oxford
United Kingdom
OX2 6GG
01865 617942
andrew.farmer@phc.ox.ac.uk

Type(s)

Public

Contact name

Miss Bethany Lawson

Contact details

Department of Primary Care Health Sciences
Radcliffe Observatory Quarter
Woodstock Road
Oxford
United Kingdom
OX2 6GG
01865 289259
bethany.lawson@phc.ox.ac.uk

Additional identifiers**Integrated Research Application System (IRAS)**

249148

Protocol serial number

PID:13866 (internal sponsor reference)

Study information**Scientific Title**

sElf-management anD support proGrammE (EDGE2) for COPD

Acronym

EDGE2

Study objectives

The aim of EDGE2 is to assess the feasibility of extracting and linking hospital data with self-monitoring data collected in a community setting by a digital health system.

Ethics approval required

Old ethics approval format

Ethics approval(s)

Approved 18/12/2018, London - Surrey Research Ethics Committee (Surrey University, London, GU2 7XH, United Kingdom; +44 (0)207 104 8088; surrey.rec@hra.nhs.uk), ref: 18/LO/1939

Primary study design

Observational

Study design

Prospective cohort study

Study type(s)

Other

Health condition(s) or problem(s) studied

Chronic Obstructive Pulmonary Disease (COPD)

Interventions

EDGE2 is a cohort study and all participants will access the intervention. The intervention will comprise an internet-linked tablet computer and two monitoring devices to support their self-management of their condition. The monitoring devices will capture oxygen saturation levels, pulse rate and step count. Patients will be encouraged to wear the physical activity monitor and to use the pulse oximeter daily for 6 months after discharge. Using the tablet computer, participants will be able to answer questions about their COPD symptoms (daily) and mood (monthly) and will be able to access educational videos about COPD.

Intervention Type

Device

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

EDGE2 Home Monitoring System

Primary outcome(s)

1. Proportion of participants in whom data can be obtained and matched from the in-hospital system and the EDGE2 platform – extract and link in-hospital data with other study data for each patient, success pre-defined as 80-90% – initiation of linkage will take place upon provision of consent with in-hospital data retrieved from point of index attendance/admission to discharge.
2. Proportion where there is sufficient data to provide clinically relevant data for use across the care pathway – obtain and link in-hospital data with other study data for each patient, success pre-defined as 80-90% – initiation of linkage will take place upon provision of consent with in-hospital data retrieved from point of index attendance/admission to discharge.

Key secondary outcome(s)

1. Quality of life - using St George's Respiratory Questionnaire for COPD (SGRQ-C) - at baseline and 6 months after hospital discharge
2. Generic health status - using the EuroQol-5 Dimension (EQ-5D-5L) scale - at baseline and 6 months after hospital discharge
3. Number of hospital admissions - at baseline and 6 months after hospital discharge
4. Number of ICU admissions - at baseline and 6 months after hospital discharge
5. Number of contacts with health professionals - at baseline and 6 months after hospital discharge
6. Use of medications - at baseline and 6 months after hospital discharge
7. Physical function – using the sit-to-stand test - at baseline, 4 weeks, 12 weeks and 24 weeks after hospital discharge
8. Death - using NHS Digital's Data Access Request Service
9. Attendance and admissions to hospital – using NHS Digital's Data Access Request Service

Completion date

27/01/2025

Eligibility

Key inclusion criteria

1. Be willing and able to give informed consent
2. Be aged 40 years or older
3. Have a clinical diagnosis of chronic obstructive pulmonary disease recorded in their medical history
4. Be a current or ex-smoker
5. Be an acute hospital attendance/admission for an exacerbation of chronic obstructive pulmonary disease or pulmonary infection
6. Be able to complete questionnaires (electronic or paper) and use the tablet computer
7. Confirm their post-discharge destination is not a medical facility or prison
8. Lives in Oxfordshire or surrounding counties
9. Be able to adequately understand verbal and written English

Participant type(s)

Patient

Healthy volunteers allowed

No

Age group

Mixed

Lower age limit

40 Years

Upper age limit

110 Years

Sex

All

Total final enrolment

41

Key exclusion criteria

1. Another significant lung disease e.g. lung cancer
2. Chronic heart failure defined by the New York Heart Association classification system as severe (Grade IV)
3. Have a life expectancy of less than six months or be on a palliative pathway

Date of first enrolment

13/05/2019

Date of final enrolment

31/03/2022

Locations**Countries of recruitment**

United Kingdom

England

Study participating centre

Oxford University Hospitals NHS Foundation Trust

John Radcliffe Hospital

Headley Way

Headington

Oxford

England

OX3 9DU

Sponsor information

Organisation

Clinical Trials and Research Governance – University of Oxford

ROR

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

Funder(s)

Funder type

Government

Funder Name

National Institute for Health Research Oxford Biomedical Research Centre

Funder Name

Engineering and Physical Sciences Research Council

Alternative Name(s)

UKRI Engineering and Physical Sciences Research Council, Engineering and Physical Sciences Research Council - UKRI, Engineering & Physical Sciences Research Council, Science Research Council, Science and Engineering Research Council, EPSRC, SRC, SERC

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location
United Kingdom

Results and Publications

Individual participant data (IPD) sharing plan

IPD sharing plan summary
Data sharing statement to be made available at a later date

Study outputs					
Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
HRA research summary	version 3	06/10/2021	28/06/2023	No	No
Other unpublished results			28/01/2026	No	No
Protocol file			16/11/2023	No	No
Study website	Study website	11/11/2025	11/11/2025	No	Yes