# Impact of receiving recorded mental health recovery narratives on quality of life in people experiencing non-psychosis mental health problems (NEON-O Trial)

Submission date 02/12/2019	<b>Recruitment status</b> No longer recruiting	[X] Prospectively registered		
		[X] Protocol		
<b>Registration date</b> 09/01/2020	<b>Overall study status</b> Completed	[X] Statistical analysis plan		
		[X] Results		
Last Edited 13/02/2024	<b>Condition category</b> Mental and Behavioural Disorders	Individual participant data		

### Plain English summary of protocol

### Background and study aims

Mental health recovery narratives are people's stories of recovery from mental health problems. Recovery narratives can be presented in a form that cannot change, such as text, audio or video, in which case we have called them "recorded" recovery narratives.

The Narrative Experiences Online (NEON) study has been investigating whether receiving recorded mental health recovery narratives can improve quality of life for people affected by mental health difficulties. NEON has identified a range of potential benefits, including feeling more hopeful or connected to others, and learning about other people's experiences. NEON is currently being trialled in individuals with experience of psychosis to investigate whether receiving access to mental health recovery narratives can provide benefits (http://www.isrctn. com/ISRCTN11152837).

Our research suggests that any benefits of receiving recovery narratives might not be specific to a mental health experience or diagnosis. We are interested in understanding whether the NEON Intervention needs to be specialised to a group of people with similar experiences (as in the NEON Trial) or whether it can be used across a broad population. In parallel with the NEON Trial, we are also conducting the NEON-O Trial, where O refers to Other. NEON-O invites participation from people with experience of any mental health problem other than psychosis.

### Who can participate?

People aged 18 years or over, who have experienced distress associated with mental health concerns in the past 6 months and have experience of mental health problem other than psychosis in the past 5 years. Participants need to be competent in English, and can use a computer or smartphone (with support if needed). Participants do not need to have received any medical support for their mental health difficulties; no diagnosis is required, and we welcome participation from people who reject the concept of diagnosis.

#### What does the study involve?

Taking part in NEON-O will involve being randomly allocated to either receive access to the

NEON Intervention, an interactive website providing access to hundreds of recovery narratives, or to receive access to the NEON Intervention after one year. Participants can use the NEON Intervention as much or as little as they want. At four points during the year, participants will need to provide some information about themselves and their experiences, using some online forms, they may also be invited to take part in an interview about their experiences, and their usage of the NEON Intervention will be logged. Results will be published in publications which are available to all.

What are the possible benefits and risks of participating?

Benefits include obtaining access to a diverse set of recovery narratives assembled by the NEON study team, and making a contribution to research which will shape clinical practice. Some recovery narratives might help participants feel more hopeful about their own future, or more connected to others with similar experiences. People can sometimes experience distress as they read, watch or listen to a recovery narrative, but this is typically short-lived. There is some evidence that encountering descriptions of self-harm in recovery narratives might contribute to recipients emulating these behaviours if they are at risk of self-harm.

Where is the study run from?

- The study is run from 12 sites.
- 1. Nottinghamshire Healthcare NHS Foundation Trust (UK)
- 2. Sussex Partnership NHS Foundation Trust (UK)
- 3. Lincolnshire Partnership NHS Foundation Trust (UK)
- 4. East London NHS Foundation Trust (UK)
- 5. South London and Maudsley NHS Foundation Trust (UK)
- 6. Derbyshire Healthcare NHS Foundation Trust (UK)
- 7. Devon Partnership NHS Foundation Trust (UK)
- 8. North East London NHS Foundation Trust (UK)
- 9. Oxford Health NHS Foundation Trust (UK)
- 10. Leicestershire Partnership NHS Trust (UK)
- 11. Cornwall Partnership NHS Foundation Trust (UK)
- 12. Somerset Partnership NHS Foundation Trust (UK)

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

The NEON-O Trial opened on 9th March 2020, and participants can currently join the trial up until 30th April 2021.

Who is funding the study?

The National Institute for Health Research (NIHR) in the United Kingdom.

Who is the main contact? Dr Stefan Rennick-Egglestone stefan.egglestone@nottingham.ac.uk

### Study website

http://www.researchintorecovery.com/neontrial

# **Contact information**

**Type(s)** Scientific **Contact name** Dr Stefan Rennick-Egglestone

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## Additional identifiers

EudraCT/CTIS number Nil known

IRAS number 249015

**ClinicalTrials.gov number** Nil known

Secondary identifying numbers IRAS Project ID: 249015

# Study information

### Scientific Title

The NEON-O Trial: a two-arm randomised controlled trial in which people with experience of nonpsychosis mental health problems receive access to (arm 1) versus receive 1 year delayed access to (arm 2) recorded mental health recovery narratives delivered online, with quality of life as a primary outcome, and hope, empowerment, meaning in life and symptomatology as secondary outcomes

Acronym NEON-O Trial

### **Study objectives**

Current study hypothesis as of 25/11/2020:

Participants receiving access to mental health recovery narratives will have a clinically-important increase in quality of life at 1-year follow-up compared with those not receiving access to mental health recovery narratives in this period.

Previous study hypothesis:

This is a feasibility trial which will develop knowledge to support the design of a future definitive trial.

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Approved 13/12/2019, Leicester Central Research Ethics Committee (The Old Chapel, Royal Standard Place, Nottingham, NG1 6FS; Tel: +44 (0)207 104 8234; Email: NRESCommittee. EastMidlands-LeicesterCentral@nhs.net), REC ref: 19/EM/0326

### Study design

Interventional randomized controlled trial with no masking delivered online with recruitment across England control of treatment as usual for one year followed by access to the intervention and 1:1 randomised allocation

### Primary study design

Interventional

### Secondary study design

Randomised controlled trial

#### Study setting(s) Internet/virtual

### Study type(s)

Other

### Participant information sheet

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

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

Mental health

### Interventions

Participants will access the stories through an online interface called the NEON Intervention. The NEON Intervention draws on stories in the NEON Collection. This has been assembled from published to stories for which we have collected consent for use, and from donations of stories directly to the study.

Equal allocation across arms.

Arm 1: treatment as usual plus access to a collection of recorded recovery stories presented online for one year.

Arm 2: treatment as usual for one year, followed by access to recorded recovery narratives

### Randomisation:

In advance of trial start, an independent statistician will generate a file of random numbers, with properties as described in our protocol. This will be uploaded to the online interface, and used to randomly allocate participants.

### Intervention Type

Other

### Primary outcome measure

Manchester Short Assessment (MANSA) of health-related quality of life, captured at 1 week, 12 weeks and 52 weeks (primary endpoint) after baseline.

### Secondary outcome measures

Secondary outcome measures will be assessed at baseline, one week, 12 weeks and 52 weeks after baseline:

- 1. Hope will be assessed through the Herth Hope Index
- 2. Meaning in Life will be assessed through the Meaning in Life Questionnaire
- 3. Empowerment will be assessed through the Mental Health Confidence Scale
- 4. Symptomatology will be assessed through the CORE-10

### Overall study start date

06/08/2018

**Completion date** 

22/09/2022

# Eligibility

### Key inclusion criteria

- 1. Aged 18 years and above
- 2. Any gender
- 3. Experinece of mental health problem other than psychosis in the last five years
- 4. Experience of mental health-related distress in previous 6 months
- 5. Resident in England
- 6. Able to understand written and spoken English
- 7. Able to access or be supported to access the internet
- 8. Able to give online informed consent.

### Participant type(s)

Patient

**Age group** Adult

Lower age limit

18 Years

**Sex** Both

**Target number of participants** 994

**Total final enrolment** 1023

Key exclusion criteria

1. If included in the NEON Trial (ISRCTN number: ISRCTN11152837)

Date of first enrolment 09/03/2020

Date of final enrolment 26/03/2021

### Locations

**Countries of recruitment** England

United Kingdom

**Study participating centre Nottinghamshire Healthcare NHS Foundation Trust** Duncan MacMillan House, Porchester Road Nottingham United Kingdom NG3 6AA

**Study participating centre Sussex Partnership NHS Foundation Trust** Swandean Arundel Road Worthing United Kingdom BN13 3EP

**Study participating centre Lincolnshire Partnership NHS Foundation Trust** St George's Long Leys Road Lincoln United Kingdom LN1 1FS

**Study participating centre East London NHS Foundation Trust** Robert Dolan House Trust Headquarters 9 Alie Street London United Kingdom E1 8DE

#### **Study participating centre South London and Maudsley NHS Foundation Trust** Bethlem Royal Hospital Monks Orchard Road Beckenham United Kingdom BR3 3BX

#### Study participating centre Derbyshire Healthcare NHS Foundation Trust

Ashbourne Centre Kingsway Hospital Kingsway Derby United Kingdom DE22 3LZ

### Study participating centre

**Devon Partnership NHS Foundation Trust** Wonford House Dryden Road Exeter United Kingdom EX2 5AF

#### Study participating centre

North East London NHS Foundation Trust The West Wing CEME Centre Marsh Way Rainham United Kingdom RM13 8GQ

### Study participating centre

**Oxford Health NHS Foundation Trust** Warneford Hospital Warneford Lane Headington Oxford United Kingdom OX3 7JX

Study participating centre

### Leicestershire Partnership NHS Trust

HQ Bridge Park Plaza Bridge Park Road Thurmaston Leicester United Kingdom LE4 8PQ

#### Study participating centre Cornwall Partnership NHS Foundation Trust Carew House Beacon Technology Park Dunmere Road Bodmin United Kingdom PL31 2QN

**Study participating centre Somerset Partnership NHS Foundation Trust** 2nd Floor, Mallard Court Express Park Bristol Road Bridgwater United Kingdom TA6 4RN

### Sponsor information

**Organisation** Nottinghamshire Healthcare NHS Foundation Trust

#### Sponsor details Duncan Macmillan House Porchester Road Mapperley Nottingham England United Kingdom NG3 6AA +44 (0)1157484321 Research@nottshc.nhs.uk

#### Sponsor type

Hospital/treatment centre

### Website

https://www.nottinghamshirehealthcare.nhs.uk/

## Funder(s)

Funder type Not defined

Funder Name National Institute for Health Research

## **Results and Publications**

### Publication and dissemination plan

The trialists will publish the trial protocol, which will incorporate a statistical analysis plan.

Dissemination through:

- 1. Publication of trial report in a journal with an international audience
- 2. Publication of a lay summary through the study website

### Intention to publish date

01/02/2024

### Individual participant data (IPD) sharing plan

Current participant level data sharing statement as of 20/10/2023:

Data access is controlled to protect the confidentiality of trial participants, and in particular to avoid re-identification through combination of multiple data files. Data will be available on reasonable request until the end of the retention period, supervised by the study sponsor. After the retention period, availability through the study sponsor or Chief Investigator may be provided at their discretion. Contact the study sponsor through Research@nottshc.nhs.uk citing IRAS ID 249015. To obtain access, an end-user license must be signed by an authorised representative. Requests can be denied if the sponsor has reason to believe that the requestor has malicious intent, and whilst research publications are being generated by the study team or investigators. Only anonymous and pseudonymous elements of the datasets used or analysed during the study will be available. Informed consent information has been retained for audit but will not be shared. Some categories of demographic data will be redacted to avoid re-identification. A data dictionary will be provided.

Previous participant level data sharing statement:

Enquiries should be addressed to m.slade@nottingham.ac.uk or to the Research and Innovation department of the study sponsor, Nottinghamshire Healthcare NHS Foundation Trust. The trialists will provide a statement clarifying data availability in their published trial protocol.

# **IPD sharing plan summary** Available on request

### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
Protocol article	protocol	20/07/2020	22/07/2020	Yes	No
Protocol article	updated protocol	29/01/2022	31/01/2022	Yes	No
<u>Statistical Analysis Plan</u>		20/05/2023	22/05/2023	Yes	No
Other publications	Development and delivery cost	07/11/2022	27/06/2023	Yes	No
Interim results article	Baseline data analysis	27/06/2023	13/07/2023	Yes	No
HRA research summary			26/07/2023		No
<u>Results article</u>		01/02/2024	13/02/2024	Yes	No