

The use of a new virtual reality software in psychiatric inpatient wards

Submission date 06/10/2025	Recruitment status Recruiting	<input type="checkbox"/> Prospectively registered
		<input type="checkbox"/> Protocol
Registration date 10/11/2025	Overall study status Ongoing	<input type="checkbox"/> Statistical analysis plan
		<input type="checkbox"/> Results
Last Edited 04/12/2025	Condition category Mental and Behavioural Disorders	<input type="checkbox"/> Individual participant data
		<input checked="" type="checkbox"/> Record updated in last year

Plain English summary of protocol

Background and study aims

The current study aims to examine the feasibility and acceptability of a virtual reality (VR) app for use on inpatient psychiatric wards. The study will be completed in two stages. Stage One is a co-design stage, and Stage Two is the Pilot stage.

Who can participate?

Adult staff who are currently working on an inpatient ward at Springfield University Hospital, and adult inpatients at Springfield University Hospital.

What does the study involve?

A VR hypnotherapy app that has been developed in collaboration with the company Phase Space will be trialled with psychiatric inpatients.

Stage One

Sixteen patients and six staff members will be recruited at this stage. The first eight patients will test the headset and provide feedback. Based on this, the app will be modified by Phase Space and then trialled by an additional eight patients. Concurrently with the recruitment of the 16 patients, six staff members from the ward will be recruited to complete an interview about the feasibility and use of VR headsets on the ward. The total feedback will be taken back to Phase Space, which will further modify the app to be implemented in the pilot stage.

Stage Two

The VR app will be piloted on fifteen inpatient participants. The full design of the pilot stage will be driven by the feedback given in Stage one. Participants will complete baseline questionnaires before the use of the headset, which will then be repeated when they have finished trialling the software. The measures will include a demographics questionnaire, as well as questionnaires regarding stress and anxiety, as well as other psychiatric symptoms. Patient participants will also be asked to complete an interview after they have finished participation to provide feedback on the use of the VR software. A semi-structured questionnaire will be completed with six staff participants, where they will be asked to report any observations they made regarding those who used the VR headset, as well as any additional feedback they have regarding the implementation of the headsets on inpatient wards.

What are the possible benefits and risks of participating?

There are no guaranteed benefits to taking part in the study. The use of the VR headset may provide participants with feelings of reduced stress. Study involvement also benefits the future of the study, as feedback will be used to develop the software further so it can possibly be implemented in clinical settings. All participants will also receive a gift voucher as a thank you for participation (£20 for Stage One, £30 for Stage Two).

It is not anticipated that any significant risks will be involved in the study. It is possible to feel cybersickness when using a VR headset. If this occurs, participants can take off the headset to let the symptoms subside. Participants can also withdraw involvement in the study at any point. It is not anticipated that participants will feel distressed due to study involvement, but if they do, the researchers can alert their care team and participants. If there are any technological problems with the VR headset, the researcher present will be trained to work through the problems.

Where is the study run from?

Springfield University Hospital, UK.

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

February 2025 to January 2026

Who is funding the study?

1. Medical Research Council (MRC), UK
2. Phase Space Ltd, UK

Who is the main contact?

Megan Cartier, megan.cartier@swlstg.nhs.uk

Contact information

Type(s)

Scientific, Principal investigator

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Additional identifiers**Central Portfolio Management System (CPMS)**

63655

Integrated Research Application System (IRAS)

351569

ClinicalTrials.gov (NCT)

NCT06917456

Protocol serial number

APP49548

Study information**Scientific Title**

The codesign and evaluation of a novel virtual reality intervention for use in psychiatric inpatient wards

Study objectives

Stage One: Gain feedback and reactions of patients and staff on the use of the VR headset to refine the VR content and provide input on the design of the Pilot

Stage Two: Explore the feasibility of the use of VR in inpatient wards

Ethics approval required

Ethics approval required

Ethics approval(s)

approved 16/07/2025, East of England - Cambridge Central Research Ethics Committee (2 Redman Place, Stratford, London, E20 1JQ, United Kingdom; +44 (0)2071048285; cambridgecentral.rec@hra.nhs.uk), ref: 25/EE/0097

Study design

Single-site interventional pilot study

Primary study design

Interventional

Study type(s)

Other

Health condition(s) or problem(s) studied

Mental health conditions

Interventions

This study examines the use of a virtual reality headset for mental health conditions.

There is no randomisation plan for the study as all participants will test out the virtual reality headset. They will use the headset to watch a seven-minute immersive experience, aimed at helping with feelings of stress and aiding in relaxation. Participants receive information on the programme before use and are aided in the use of the headset by two trained researchers. The study of the headset will be done in-person within the wards within the sensory or calm rooms, so participants have a quiet space to try out the headset.

Intervention Type

Device

Phase

Not Applicable

Drug/device/biological/vaccine name(s)

Virtual reality hypnotherapy app and headset

Primary outcome(s)

1. Stage One: Gather feedback and reactions of both staff and patients on the use of the VR headset to refine the content and provide input on the design of the Pilot, measured using qualitative interviews with patients and staff at the end of study involvement
2. Stage Two: Explore the feasibility of the use of VR in inpatient wards, measured using qualitative interviews with patients and staff at the end of study involvement

Key secondary outcome(s))

1. Stage One: Produce an implementation model to be tested in a Pilot, identify any areas requiring further modification, learn how to prepare staff and patients for the VR experience, identify the best location for the VR experience, and understand the practicalities of the headsets, measured using qualitative interviews with staff and patients at the end of study involvement
2. Stage Two: Explore the impact of the VR on patients measured using the Simulator Sickness Questionnaire, Patient Health Questionnaire-8 (PHQ-8), Generalized Anxiety Disorder 7-item Scale (GAD-7), Dissociative Experiences Scale – II (DES-II), Brief Psychiatric Rating Scale (BPRS), Positive and Negative Syndrome Scale (PANSS) at baseline and at the end of study involvement

Completion date

31/01/2026

Eligibility

Key inclusion criteria

Staff:

1. Currently working on an inpatient ward at Springfield University Hospital
2. Able to provide informed consent

Patients:

1. Aged 18-65 years old
2. Current inpatient at Springfield University Hospital
3. Able to provide informed consent

Participant type(s)

Health professional, Patient

Healthy volunteers allowed

No

Age group

Mixed

Lower age limit

18 years

Upper age limit

65 years

Sex

All

Total final enrolment

0

Key exclusion criteria

Staff:

1. Not currently working on an inpatient ward at Springfield University Hospital
2. Unable to provide informed consent

Patients:

1. Unable to provide informed consent
2. Has a history of seizures
3. Has a pacemaker

Date of first enrolment

13/08/2025

Date of final enrolment

31/01/2026

Locations

Countries of recruitment

United Kingdom

England

Study participating centre

South West London & St. George's Mental Health NHS Trust

Springfield University Hospital

Trinity Building

15 Springfield Drive

London

England

SW17 0YF

Sponsor information

Organisation

South West London and St George's Mental Health NHS Trust

ROR

<https://ror.org/003pb1s55>

Funder(s)

Funder type

Government

Funder Name

Medical Research Council

Alternative Name(s)

Medical Research Council (United Kingdom), UK Medical Research Council, MRC

Funding Body Type

Government organisation

Funding Body Subtype

National government

Location

United Kingdom

Funder Name

Phase Space

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?
Participant information sheet	Participant information sheet	11/11/2025	11/11/2025	No	Yes