# Can a brief relational talking therapy help adults who self-harm?

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
18/10/2022		[X] Protocol		
Registration date	Overall study status Completed	Statistical analysis plan		
21/10/2022		[X] Results		
Last Edited	Condition category	Individual participant data		
11/12/2024	Mental and Behavioural Disorders			

#### Plain English summary of protocol

Background and study aims

Self-harm refers to when a person intentionally harms themselves, for example by cutting themselves or taking an overdose. Self-harm can occur both with and without suicidal intent. Research suggests that in the UK rates of self-harm have increased. It is a major health concern because self-harm is associated with a greater risk of suicide and emotional distress. Evidence of which therapies are most helpful for people who self-harm is limited. Some therapies that are currently used are also very intensive (e.g. Dialectical Behaviour Therapy), can be difficult to access, and do not suit everybody. Cognitive Analytic Therapy (CAT) is a talking therapy that has the potential to help people who self-harm. It may provide a helpful alternative for people struggling with self-harm that may not be eligible for or able to access other therapies. Before a large-scale clinical trial of CAT for self-harm can be undertaken, it is helpful to conduct a smaller-scale feasibility trial. The aim of the current study is therefore to undertake a feasibility study of CAT for adults who self-harm. Data will be collected to help answer important feasibility uncertainties (e.g. can we recruit an adequate sample).

#### Who can participate?

Adults with experiences of self-harm from NHS mental health services, including IAPT (Improving Access to Psychological Therapy) services and other community-based mental health services

#### What does the study involve?

Participants will first be asked to complete a baseline assessment with a researcher, answering questions about their thoughts, feelings, and recent difficulties. Participants will then be randomly put into one of two groups. One group will receive the therapy, CAT, plus the usual treatment. The other group will just have access to their usual treatment. There will be further meetings with the researcher to do more assessments at 12 and 18 weeks after being put into the group. Some participants will also be invited to take part in a more in-depth interview or complete additional questionnaires online, but these parts of the study are optional. We are hoping to recruit 60 people to take part overall.

What are the possible benefits and risks of participating? We do not yet know how helpful CAT will be for people who self-harm, which is why we are doing this research. Taking part in research can be an interesting and rewarding experience though. Taking part in this study will help us better understand which treatments might help people who self-harm. Taking part may therefore feel like a positive experience for some people and a way to help others. The study will involve being asked questions about difficult experiences like self-harm, and this may be uncomfortable to distressing for some people. Participants will not have to answer any questions they do not wish to and will be able to leave the study at any time. The researchers will be trained to respond to distress and will be able to offer breaks during meetings and provide advice on further sources of support.

#### Where is the study run from?

The lead site is the study is Greater Manchester Mental Health and Social Care NHS Foundation Trust. The second site is at Rotherham, Doncaster, and South Humber NHS Foundation Trust

When is the study starting and how long is it expected to run for? December 2022 to December 2024

Who is funding the study? National Institute for Health and Care Research (NIHR) (UK)

Who is the main contact?

Dr Peter Taylor (Project lead), peter.taylor-2@manchester.ac.uk

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### **Contact information**

#### Type(s)

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#### Additional identifiers

Clinical Trials Information System (CTIS)

Nil known

Integrated Research Application System (IRAS)

318068

ClinicalTrials.gov (NCT)

Nil known

Protocol serial number

CPMS 54183, IRAS 318068

## Study information

#### Scientific Title

Relational Approach to Treating Self-Harm (RelATe): a feasibility study of cognitive analytic therapy for people who self-harm

#### Acronym

RelATe

#### **Study objectives**

As this is a feasibility trial we have progression criteria rather than hypotheses. These are outlined below.

A traffic light system (Green = progress to definitive trial; Amber = modification needed; Red = do not progress) has been adopted to guide the decision to progress:

- 1. Recruitment rates: Data on the ability to randomise 60 participants within 12 months (5 participants per month). Green: ≥80%. Amber: 60-79%. Red: ≤59%.
- 2. Retention rate (completion of end-of-treatment follow-up assessments). Green: ≥80%. Amber 60-79%. Red ≤59%.
- 3. Missing data on primary outcomes at assessment. Green: <15%. Amber: 16-25%. Red >25%.
- 4. Adherence to treatment: Percentage of participants receiving the minimum dose of therapy (≥4 sessions) within a 10-week treatment window. Green: ≥80%. Amber: 60-79%. Red: ≤59%.

#### Ethics approval required

#### Old ethics approval format

#### Ethics approval(s)

Approved 07/11/2022, Greater Manchester West Research Ethics Committee (Barlow House, 3rd Floor, HRA NRES Centre Manchester, M1 3DZ, UK; +44 (0)207 104 8379; gmwest.rec@hra.nhs. uk), ref: 22/NW/0317

#### Study design

Randomized interventional study

#### Primary study design

Interventional

#### Study type(s)

Treatment

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

Mental health

#### Interventions

Cognitive Analytic Therapy (CAT) will be delivered by a qualified band 7 or 8 practitioner with appropriate core professional training (e.g., clinical psychologist) who has undertaken additional training in CAT (completing at least the first year of post-qualification CAT practitioner training). Therapists will be supervised by an accredited CAT therapist on a fortnightly basis. The therapy will follow a standard eight-session CAT model, with a specific focus on self-harm as the presenting problem. Sessions 1-3 would typically focus on developing a shared understanding of difficulties and identifying unhelpful patterns that maintain problems, using visual (drawn) mapping of experiences and therapeutic letters to assist in this process. Sessions 4-8 would typically focus on the recognition of unhelpful patterns identified so far, and the use of alternative, more helpful ways of responding to experiences and challenges, referred to as "exits". A further follow-up session will take place up to 8 weeks after the end of therapy, based on recommendations from Patient and Public Involvement (PPI) meetings. Therapy sessions will last 50 to 60 minutes and typically occur weekly. Based on recommendations from PPI contributors, sessions could be undertaken either in person or remotely via a video call.

#### Intervention Type

Other

#### Primary outcome(s)

Measured using patient records:

- 1. Recruitment rates will be recorded across the 12 month recruitment period
- 2. Participant retention rate (completion of end-of-treatment follow-up assessments) will be recorded over 18 weeks (the length of time in the study)
- 3. Participant missing data on the primary outcomes (self-harm behaviour and urges) at assessment will be recorded over 18 weeks (the length of time in the study)
- 4. Adherence to treatment will be recorded as the percentage of participants receiving the minimum dose of therapy (≥4 sessions) within the 10-week treatment window.

#### Key secondary outcome(s))

1. Self-harm measured with the Self-Injurious Thoughts and Behaviours Interview-Short Form (SITBI) at baseline, 12 weeks, and 18 weeks

- 2. Self-harm urges measured with the Alexian Brothers Urges to Self-Injure Scale (ABUSI) at baseline, 12 weeks, and 18 weeks
- 3. Dependence on self-ham measured with the experiences of Self-Injury Questionnaire Positive Beliefs subscale at baseline, 12 weeks, and 18 weeks
- 4. Personality structure measured with the Personality Structure Questionnaire at baseline, 12 weeks, and 18 weeks
- 5. Emotional distress measured with the Kessler Distress Scale at baseline, 12 weeks, and 18 weeks
- 6. Interpersonal problems measured with the Inventory of Interpersonal Problems-32 at baseline, 12 weeks, and 18 weeks
- 7. Health status measured with the EQ-5D-5L at baseline, 12 weeks, and 18 weeks
- 8. Adverse experiences from therapy measured with the Adverse Experiences in Psychotherapy scale (AEP) at 12 weeks

#### Completion date

11/12/2024

## **Eligibility**

#### Key inclusion criteria

- 1. Aged 18 years or older
- 2. Three or more episodes of self-harm in the past year, confirmed via the Self-Injurious Thoughts and Behaviours Interview
- 3. Can be safely seen in an outpatient clinical context in which treatment is being provided as judged by their clinical team or referrer

#### Participant type(s)

**Patient** 

#### Healthy volunteers allowed

No

#### Age group

Adult

#### Lower age limit

18 years

#### Sex

All

#### Total final enrolment

60

#### Key exclusion criteria

- 1. Participants will be excluded if they have a moderate-to-severe intellectual disability (IQ:< 70) affecting their eligibility for community mental health services as judged by their clinical team or referrer
- 2. Organic cerebral disease/injury affecting receptive and expressive language comprehension as judged by their clinical team

- 3. Non-English speaking to the degree that the participant is unable to answer questions and give written informed consent
- 4. Imminent and immediate risk to self or others, operationalised as the presence of active suicidal intent or planning to end one's life in the near future (e.g. next week). Where individuals are excluded on this basis, with the person's consent, the researcher will aim to recontact them and the referrer in approximately one month's time (or a time period agreed upon in collaboration with the individual) to determine if the risk has subsided to a point where they are now eligible.
- 5. Currently an inpatient
- 6. Experiencing a current, active episode of psychosis or mania
- 7. Currently receiving another active one-to-one psychological therapy

### Date of first enrolment

13/02/2023

Date of final enrolment 29/02/2024

#### Locations

#### Countries of recruitment

United Kingdom

England

## Study participating centre Prestwich Hospital

Bury New Road Prestwich Manchester United Kingdom M25 3BL

## Study participating centre Rotherham Doncaster and South Humber NHS Foundation Trust

Woodfield House Tickhill Road Doncaster United Kingdom DN4 8QN

## Sponsor information

Greater Manchester Mental Health NHS Foundation Trust

#### **ROR**

https://ror.org/05sb89p83

## Funder(s)

#### Funder type

Government

#### Funder Name

National Institute for Health and Care Research Central Commissioning Facility (CCF); Grant Codes: NIHR203515

#### 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

#### **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. The decision on which repository to use has not yet been made but will be reviewed as the trial progresses. Qualitative data will not be shared due to the greater challenges in fully anonymising such data. Data will be made available to other researchers to access once the main trial results have been published. Participants will be informed that data may be shared in this way.

#### IPD sharing plan summary

Stored in publicly available repository

#### Study outputs

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
<u>Protocol article</u>		18/07/2024	19/07/2024	Yes	No
HRA research summary			28/06/2023	No	No
Other unpublished results		11/12/2024	11/12/2024	No	No