# Psychological treatment for men with intellectual and/or developmental disabilities and harmful sexual behaviour

Submission date	Recruitment status No longer recruiting	[X] Prospectively registered		
11/05/2021		☐ Protocol		
Registration date	Overall study status	Statistical analysis plan		
21/05/2021	Ongoing	☐ Results		
Last Edited	Condition category	Individual participant data		
20/11/2023	Mental and Behavioural Disorders	Record updated in last year		

#### Plain English summary of protocol

Background and study aims

There is considerable public concern about men who commit sexual offences. Such men are typically offered group cognitive behaviour therapy (CBT), both in prison and in the community. This form of therapy for sex offenders has been repeatedly evaluated and most trials have found it to be effective in reducing reoffending. However, men who show harmful sexual behaviours (HSB) and have intellectual and or developmental disabilities (IDD) have often been excluded from this form of treatment so that in the past they have rarely been offered CBT.

This proposed research project aims to evaluate the effectiveness of the SOTSEC-ID CBT group for men with IDD and harmful sexual behaviour, using the gold standard method for evaluating the effectiveness of treatments, i.e. a randomised controlled trial.

Who can participate?

Men with intellectual and/or developmental disabilities and harmful sexual behaviour

What does the study involve?

The plan is for groups of men with IDD who show harmful sexual behaviours to be assessed for sexual knowledge, victim empathy, cognitive distortions, and harmful sexual behaviours, and then the groups will be allocated randomly to be treated with SOTSEC-ID group CBT (with risk management) or TAU (treatment as usual). The men will be re-assessed at the end of six months and followed up at 1 year (all groups) and 2 years(intervention groups only).

What are the possible benefits and risks of participating?

Possible benefits include improvements (reductions) in harmful sexual behaviours, and cognitive distortions, and improvements in sexual knowledge and empathy. Risks include that this is lengthy treatment and intensive which may not suit everyone.

Where is the study run from? University of Kent (UK)

When is the study starting and how long is it expected to run for? From October 2021 to September 2025

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

Who is the main contact? Prof Glynis Murphy, g.h.murphy@kent.ac.uk

#### Study website

https://research.kent.ac.uk/safer-idd/research/the-new-hasb-idd-trial/

#### Contact information

#### Type(s)

Scientific

#### Contact name

Prof Glynis Murphy

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#### Additional identifiers

#### **EudraCT/CTIS** number

Nil known

#### **IRAS** number

291027

#### ClinicalTrials.gov number

Nil known

#### Secondary identifying numbers

IRAS 291027

## Study information

Scientific Title

RCT of group CBT for men with intellectual and/or developmental disabilities and harmful sexual behaviour (HaSB-IDD)

#### Acronym

HaSB-IDD

#### **Study objectives**

- 1. To determine whether the SOTSEC-ID group CBT program, combined with risk management:
- 1.1. Reduces cognitive distortions in men with intellectual and/or developmental disabilities and harmful sexual behaviour
- 1.2. Prevents or reduces their further harmful sexual behavior
- 1.3. Improves their sexual knowledge, empathy, locus of control, and self-esteem, in comparison to men in the control group receiving Treatment As Usual (TAU)
- 2. To examine the costs and cost-effectiveness of this treatment
- 3. To examine therapist, carer and service user views of treatment (through smaller qualitative studies)

#### Ethics approval required

Old ethics approval format

#### Ethics approval(s)

Approved 05/01/2022, Nottingham 2 Research Ethics Committee (Equinox House, City Link, Nottingham, NG2 4LA; +44 (0)207 104 8169; nottingham2.rec@hra.nhs.uk), ref: 21/EM/0270

#### Study design

Multi-centre single-blinded cluster randomized controlled trial

#### Primary study design

Interventional

#### Secondary study design

Cluster randomised trial

#### Study setting(s)

Community

#### Study type(s)

Treatment

#### Participant information sheet

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

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

Harmful sexual behaviour in men with intellectual and/or developmental disabilities

#### **Interventions**

This is a cluster randomized trial of a form of group cognitive behavioural treatment know as SOTSEC-ID (see https://www.kent.ac.uk/tizard/sotsec). After assessing site eligibility, sites will be randomised to Arm A or Arm B. Arm A consists of 6 months of CBT treatment (using SOTSEC-ID), plus risk management. Arm B is treatment as usual (mainly risk management). Therapists

cannot be blind to treatment. Research workers collecting assessment and outcome data will be blind to treatment.

#### Intervention Type

Behavioural

#### Primary outcome measure

Cognitive distortions measured using the Questionnaire on Attitudes Consistent with Sexual Offending (QACSO) score at baseline, 6, 12, and 24 months

#### Secondary outcome measures

- 1. Harmful sexual behaviour measured using all reports from case files, carers, police, etc at baseline, 6, 12, and 24 months
- 2. Sexual knowledge measured using the General Sexual Knowledge Questionnaire at baseline, 6, 12, and 24 months
- 3. Victim empathy measured using the Victim Empathy Scale (VES-A), adapted from Beckett and Fisher's Victim Empathy Scale at baseline, 6, 12, and 24 months
- 4. Locus of control measured using the Nowicki-Strickland Locus of Control Scale at baseline, 6, 12, and 24 months
- 5. Self-esteem measured using the Rosenberg self esteem scale at baseline, 6, 12, and 24 months
- 6. Service use measured using the Client Services Receipt Inventory at baseline, 6, 12, and 24 months
- 7. Quality of life measured using the EuroQol 5-dimension 5-level (EQ-5D-5L) questionnaire at baseline, 6, 12, and 24 months

#### Overall study start date

01/10/2021

#### Completion date

30/09/2025

# **Eligibility**

#### Key inclusion criteria

- 1. Borderline or mild intellectual disability (ie an IQ below 79) and deficits in adaptive behaviours
- 2. Documented history of one or more incidents of harmful sexual behavior (HSB)
- 3. Relatively good verbal comprehension (to be judged by clinicians)
- 4. Capacity to make a decision as to whether they wish to take part in trial
- 5. Autism, additional mental health needs (as long as this would not prevent participation in the CBT group as judged by clinicians), and criminal convictions for HSB will not be reasons for exclusion

#### Participant type(s)

Mixed

#### Age group

Adult

#### Sex

Male

#### Target number of participants

240 (30 clusters with 8 men each)

#### Key exclusion criteria

- 1. Insufficient or receptive language to take part in group CBT
- 2. Mental health difficulties that would prevent him from taking part in group CBT (as judged by clinicians)
- 3. Resident in prison or in high secure services, or on probation
- 4. Does not have the capacity to make a decision as to whether to take part in the trial

#### Date of first enrolment

01/04/2022

#### Date of final enrolment

01/04/2024

#### Locations

#### Countries of recruitment

England

**United Kingdom** 

#### Study participating centre

#### Kent and Medway Partnership NHS Trust

Tarentfort Centre and Brookfield Centre Bow Arrow Lane Dartford United Kingdom DA2 6PB

#### Study participating centre

#### Northumberland Tyne and Wear NHS Foundation Trust

Rosewood Hopewood Park Hospital Ryhope Sunderland United Kingdom SR2 0NB

# Study participating centre Oxleas NHS Foundation Trust

Pinewood House Pinewood Place Dartford United Kingdom DA2 7WG

# Study participating centre Black Country Healthcare NHS Foundation Trust

Delta House Delta Point Greets Green Road West Bromwich West Bromwich United Kingdom B70 9PL

#### Study participating centre Northampton NHS Foundation Trust

Community Team for People with Learning Disabilities Second Floor Newland House Campbell Square Northampton United Kingdom NN1 3EB

# Study participating centre Nottinghamshire NHS Healthcare Trust

Low Secure & Community Forensic Directorate Wells Road Centre Mapperley Nottingham United Kingdom NG3 3AA

## Sponsor information

#### Organisation

University of Kent

#### Sponsor details

Canterbury Canterbury England United Kingdom CT2 7NS +44 (0)1227 764000 researchculture@kent.ac.uk

#### Sponsor type

University/education

#### Website

www.kent.ac.uk

#### **ROR**

https://ror.org/00xkeyj56

## Funder(s)

#### Funder type

Government

#### **Funder Name**

National Institute for Health Research

#### 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**

#### Publication and dissemination plan

The study protocol will be published in a peer-review publication. The Statical Analysis Plan will be approved by the Trial Management Group and available from the Chief Investigator post analysis. The results of the study will be disseminated via:

- 1. An overall project report to the HTA
- 2. Six open-access publications: one on the protocol; two quantitative reports of the results; two qualitative reports; one health economics paper
- 3. Two papers in professional journals (e.g. the Tizard Learning Disability Review)

- 4. A website dedicated to the project
- 5. Annual 'progress reports' to the Seattle Club Conference on Research in Intellectual and Developmental Disabilities, which is the major UK research meeting in this area
- 6. End of project presentations to the International Association for the Scientific Study of Intellectual Disability (IASSID), National Organisation for the Treatment of Abusers (NOTA), BPS forensic conference and the British Association for Behavioural and Cognitive Psychotherapy (BABCP)
- 7. Dissemination to public bodies and charities e.g. Mencap, BILD, and CBF

#### Intention to publish date

01/04/2027

#### Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are/will be available upon request from Prof Murphy, Chief Investigator (g.h.murphy@kent.ac.uk) and Prof Lee Shepstone Lead Statistician (l.shepstone@uea.ac.uk). All data will be potentially available, all of it will be already anonymised. These will not be available until October 2027.

#### IPD sharing plan summary

Available on request

#### **Study outputs**

Output type	Details	Date created	Date added	Peer reviewed?	Patient-facing?
HRA research summary			28/06/2023	No	No