# Four-day intensive treatment for obsessivecompulsive disorder - a feasibility study

<b>Submission date</b> 09/06/2022	<b>Recruitment status</b> No longer recruiting	Prospectively registered
		☐ Protocol
Registration date	Overall study status	Statistical analysis plan
10/06/2022	Completed	☐ Results
Last Edited	Condition category	Individual participant data
10/06/2022	Mental and Behavioural Disorders	<ul><li>Record updated in last year</li></ul>

### Plain English summary of protocol

Background and study aims

Obsessive compulsive disorder (OCD) is a common mental health condition where a person has obsessive thoughts and compulsive behaviours. OCD can affect men, women and children. Some people start having symptoms early, often around puberty, but it usually starts during early adulthood.

Bergen 4 Day Treatment (B4DT) is a concentrated exposure treatment (cET) developed to treat OCD. It has proven to be highly acceptable and effective in other settings. In this feasibility study we investigate the implementation, feasibility, and preliminary outcomes of B4DT treatment in HUS psychiatry settings in 5-7 therapy groups.

### Who can participate?

Adult patients who have been diagnosed with obsessive compulsive disorder.

### What does the study involve?

The study involves a pre-screening interview, the actual intensive treatment period in four consecutive days in HUS outpatient clinic within a group of 3-6 patients and at least the same number of therapists, and a follow-up at 10 days and at 3 months after the treatment.

### What are the possible benefits and risks of participating?

Based on previous research results, we assume the Bergen 4 Day Treatment model will be an effective and safe treatment that is well suited for the Finnish treatment environment. Participating in the study may be beneficial for you and ameliorate your OCD symptoms. As the exact same treatment has been in use in other Scandinavian countries for some years and shown to be safe and effective, it is unlikely to have any unexpected adverse events.

Where is the study run from? Helsinki University Hospital (Finland)

When is the study starting and how long is it expected to run for? November 2021 to April 2023 Who is funding the study? Helsinki University Hospital (Finland)

Who is the main contact? Suoma Saarni, PhD, suoma.saarni@hus.fi

### Contact information

### Type(s)

Principal investigator

#### Contact name

Dr Suoma Saarni

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### Additional identifiers

### Clinical Trials Information System (CTIS)

Nil known

### ClinicalTrials.gov (NCT)

Nil known

### Protocol serial number

Nil known

# Study information

#### Scientific Title

Bergen 4 Day Treatment (B4DT) for Obsessive-Compulsive Disorder (OCD) – an observational feasibility study

#### Acronym

B4DT-HKI-I

### **Study objectives**

We investigate the implementation, feasibility, and preliminary outcomes of B4DT treatment in HUS psychiatry settings in Finland

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Approved 06/04/2022, Helsinki and Uusimaa Hospital District (HUS) Ethics Committee (PO BOX 705, 00029, Finland; +358 40 617 5386; mirka.salin@hus.fi), ref: HUS/1085/2022

### Study design

Observational feasibility study

### Primary study design

Observational

### Study type(s)

**Treatment** 

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

Obsessive-compulsive disorder

#### **Interventions**

The patients will receive a concentrated exposure treatment (cET) conducted in four consecutive days in HUS psychiatry outpatient clinic according to Bergen four day treatment for OCD (B4DT) manual. After ten days and three weeks after treatment, patients are offered a short follow-up session to discuss their experiences during and after the active treatment period.

### Intervention Type

Behavioural

### Primary outcome(s)

Within-individual decline in Y-BOCS score before the intervention vs. after the treatment and at three months.

### Key secondary outcome(s))

Measured before the intervention vs. after the treatment and at three months.

- 1. OCD symptoms measured with OCI-R and DOCS-SF
- 2. Depression (PHQ-9)
- 3. Anxiety (GAD-7)
- 4. Sleeping problems (BIS)
- 5. Functionality (WSAS)
- 6. Well-being (WEMWBS)
- 7. Client satisfaction (CSO-8)

### Completion date

20/04/2023

# Eligibility

### Key inclusion criteria

DSM-5 Obsessive-compulsive disorder

### Participant type(s)

**Patient** 

### Healthy volunteers allowed

No

### Age group

Adult

#### Sex

Male

### Key exclusion criteria

- 1. Concurrent psychotic disorder
- 2. Substance use disorder
- 3. Imminent risk of suicide
- 4. Congenital disability
- 5. New antidepressant or antipsychotic pharmacotherapy trial started within the last six weeks
- 6. Use of benzodiazepines is allowed only for insomnia
- 7. Ongoing other evidence-based psychotherapy for OCD

### Date of first enrolment

01/05/2022

### Date of final enrolment

31/12/2022

### Locations

### Countries of recruitment

Finland

### Study participating centre Helsinki University Hospital

HUS Brain Center Department of Psychiatry P.o.Box 590 Helsinki Finland 00029 HUS

# **Sponsor information**

### Organisation

Helsinki University Hospital

#### ROR

https://ror.org/02e8hzf44

# Funder(s)

### Funder type

Hospital/treatment centre

#### Funder Name

Helsinki University Hospital

### **Results and Publications**

### Individual participant data (IPD) sharing plan

The data will be held by the Helsinki and Uusimaa Hospital District, and it will not be available because of limitations caused by the Finnish data protection legislation, content of consent requested, and research permit.

### 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