

# Effectiveness of group-delivered therapy for insomnia

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| <b>Submission date</b><br>13/10/2022   | <b>Recruitment status</b><br>No longer recruiting    | <input type="checkbox"/> Prospectively registered<br><input checked="" type="checkbox"/> Protocol            |
| <b>Registration date</b><br>17/11/2022 | <b>Overall study status</b><br>Completed             | <input checked="" type="checkbox"/> Statistical analysis plan<br><input checked="" type="checkbox"/> Results |
| <b>Last Edited</b><br>14/04/2025       | <b>Condition category</b><br>Nervous System Diseases | <input type="checkbox"/> Individual participant data   |

## Plain English summary of protocol

### Background and study aims

Insomnia is the most common sleep disorder in the general population and in clinical practice. It has been linked with an increased risk of adverse health outcomes such as cardiovascular disease, type 2 diabetes, chronic multisite pain, depression and hypertension. However, treatment options for insomnia are limited. Currently, the most common way of treating insomnia is with pharmacotherapy. Pharmacotherapy has been shown to have low-to-moderate effects and is mainly recommended for short-term use, as symptoms often return after treatment. It is therefore recommended to use cognitive behavioral therapy for insomnia (CBT-I) as first-line treatment, due to its long-term effectiveness. However, CBT-I is not widely available due to the lack of trained therapists and long waiting lists in specialist sleep clinics. Since there are limited resources for high-quality individual CBT-I, a group-delivered course for individuals with insomnia symptoms was designed by the Norwegian Directorate of Health. The course is based on core CBT-I principles including sleep hygiene, stimulus control, sleep restriction, cognitive therapy, and relaxation training. An advantage of this course is that it can be delivered to up to 15 individuals at the same time, and may thus complement traditional treatments and relieve pressure on primary care. Although this therapy has already been implemented in several municipalities in Norway, its effectiveness has never been evaluated.

The study aims to test the effectiveness of group-delivered cognitive behavioral therapy for insomnia (CBT-I) in primary healthcare. We will compare participants who received the therapy (intervention group), and those who did not (i.e., waiting list control group). The primary aim is to examine the differential change in insomnia severity at 3 months post-randomization.

Secondary aims include:

1. Examine differential changes in health-related quality of life, fatigue, depression/anxiety, and sleep diary data at 3 months post-randomization, and 6 months post-randomization.
2. Examine whether potential treatment moderators (chronotype, reactivity to stress, recruitment location, duration of insomnia, length of insomnia treatment, lifestyle etc.) influence the effectiveness of the group-delivered CBT-I in primary care.
3. Collect national registry data at 1 and 2 years post-randomization to compare sick leave days, use of prescribed medication (psychotropic medications, sedatives) and healthcare resource utilization.
4. Explore mediating analyses to identify mechanisms of change in the primary and secondary

outcomes, focusing on psychological measures of beliefs about sleep and sleep-related self-efficacy.

5. Carry out a process evaluation to assess facilitators and barriers for treatment adherence and qualitative studies to explore participants' experience of group-delivered CBT-I.

Who can participate?

Adults (18 + years old) with symptoms of insomnia (Insomnia Severity Scale of 12 or higher)

What does the study involve?

The study involves either participating in the group-delivered therapy for insomnia (intervention group) or being on a waiting list for 6 months (control group). All participants fill out a questionnaire package and a 7-day sleep diary at 4 measurement time points (weeks 1 and 4, and months 3 and 6).

What are the possible benefits and risks of participating?

For the individual participant, participation in the study provides a good opportunity to survey and improve sleep quality. Overall, the intervention is currently being offered in many Norwegian municipalities, but its effectiveness has not yet been evaluated. Evaluating the effectiveness of the intervention will strengthen the treatment of insomnia in primary care. In addition, data from the project will be used to further develop the intervention in collaboration with the Directorate of Health and sleep experts in Norway.

By taking part in the study, the participants devote time and attention to questions about health and illness, which can cause unnecessary worries. Nevertheless, these are people who already have a sleep problem and wish to treat it, and we do not consider this a major risk. Moreover, certain participants may feel concerned that their sleep and health are being carefully surveyed during the duration of the study. It is specified that all data is anonymous and that no revealing or private data will be used. Participants who are randomized to the control group end up on a waiting list for the intervention and have to wait up to 6 months before they are offered to participate in a sleep course.

Where is the study run from?

Norwegian University of Science and Technology (Norway)

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

January 2022 to December 2025

Who is funding the study?

Norwegian Foundation for Health and Rehabilitation (ExtraStiftelsen, Stiftelsen Dam & Dam Foundation) (Norway)

Who is the main contact?

Maria Hrozanova (Principal Investigator) (Norway)  
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## Contact information

**Type(s)**

Principal investigator

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**Additional identifiers**

## Clinical Trials Information System (CTIS)

Nil known

### Protocol serial number

2021/FOS374023

## Study information

### Scientific Title

Group-delivered cognitive behavioral therapy for insomnia in primary care: A pragmatic, multicenter randomized controlled trial

### Study objectives

Participants in the intervention group, who receive 4 sessions of group-delivered cognitive behavioral therapy for insomnia will experience a greater reduction of insomnia symptoms and improved health-related quality of life than the wait-list control group.

### Ethics approval required

Old ethics approval format

### Ethics approval(s)

Approved 21/06/2022, Regional Ethics Committee in Mid-Norway (NTNU/REK midt, Det medisinske fakultet, Postboks 8905, 7491 Trondheim, Norway; +4773 59 75 11; rek-midt@mh.ntnu.no), ref: 46524

### Study design

Multicenter interventional unblinded pragmatic randomized control trial

### Primary study design

Interventional

### Study type(s)

Treatment

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

Insomnia

### Interventions

Current interventions as of 13/12/2023:

Due to natural capacity limitations, some intervention group participants may have to wait to begin treatment. Waiting times over 2-3 months rarely occur. The last self-reported measurements are expected to be sent out by December 31st 2023. Thus, the data collection is expected to be finalized by January 15th 2024 at the latest.

All participants are self-referred to the study. Recruitment occurs primarily on social media (advertisements on Facebook) and through the Healthy Life Centres participating in this trial, and secondarily through information posters displayed in relevant places (e.g., GP or physiotherapy offices). Each adult interested in participating fills out a screening questionnaire, which can be found on the trial website [www.sovnbehandling.no](http://www.sovnbehandling.no). Participants who fulfil the criteria for inclusion are sent a digital consent form with detailed information about trial

participation. Those who return the signed consent form are randomized in the centre they participate. There are two trial arms:

Intervention group – participation in 4 sessions of group-delivered CBT-I. Group sizes vary from 5-15 depending on the capacity of the Healthy Life Centre. Employees deliver the intervention at the participating Healthy Life Centres (mostly physiotherapists). The intervention has been developed by the Norwegian Directorate of Health, in collaboration with the National competence centre for sleep disorders, the Public health institute and the Norwegian association for cognitive therapy. The intervention group is actively followed up for 6 months (self-reported questionnaires), and after that for 2 years (register data).

Control group – waiting list of 6 months. During the waiting period, participants fill out the same self-reported questionnaires as the intervention group. They are not offered any treatment during the 6-month waiting period. They are not prohibited from taking sleep medication at this time or seeking other forms of treatment (whether they received any treatment during their participation will be investigated at the 6-month assessment point). Control group participants are contacted once, upon randomization, by the Healthy Life Centre in their municipality, to offer them the possibility to participate in the intervention after the 6-month waiting period is finalized.

#### Previous interventions:

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#### **Intervention Type**

Behavioural

#### **Primary outcome(s)**

Insomnia measured using the Insomnia Severity Index at baseline, 4 weeks, 3 months (primary measurement point), and 6 months

#### **Key secondary outcome(s)**

1. At baseline, 4 weeks, 3 months (primary measurement point), and 6 months, all participants fill out the following secondary outcome measures:

- 1.1. Fatigue measured using the Chalder Fatigue Scale
- 1.2. Subjective sleep patterns measured using a 7-day digital sleep diary
- 1.3. Health-related quality of life measured using EuroQol EQ5D-5L
- 1.4. Mental distress measured using the Hopkins Check List
- 1.5. Chronotype measured using Brief Horne-Østberg Morningness-Eveningness questionnaire
- 1.6. Patient beliefs and attitudes about sleep measured using Dysfunctional Beliefs and Attitudes about Sleep questionnaire
- 1.7. Sleep reactivity measured using Ford Insomnia Response to Stress Test

2. At baseline, intervention group participants fill out the following secondary outcome measures:

- 2.1. Motivation for treatment measured using Nijmegen Motivation List 2
- 2.2. Treatment expectancy measured using Credibility/Expectancy Questionnaire

3. At 3 months, intervention group participants fill out the following secondary outcome measures:

- 3.1. Utility of CBT-I measured using Treatment Components Adherence Scale

4. Participant demographic information measured using a questionnaire at baseline

5. At 12 months and 24 months the following outcomes will be measured:

- 5.1. Short - and medium-term impact of group-delivered CBT-I on rates of sick leave, and medication and health resource utilization measured by accessing national health registry data
- 5.2. Overview of health service utilization measured by obtaining data from the Norwegian Patient Registry
- 5.3. Overview of prescribed medication measured by obtaining data from the Norwegian Prescription Database
- 5.4. Information on sick leave measured by obtaining data from the National Insurance Administration

### **Completion date**

31/12/2025

## **Eligibility**

### **Key inclusion criteria**

Inclusion criteria based on self-report and national registries:

1. Adults older than 18 years of age
2. Self-referred to the study
3. Insomnia Severity Index score of 12 or higher

### **Participant type(s)**

Patient

### **Healthy volunteers allowed**

No

### **Age group**

Adult

**Lower age limit**

18 years

**Sex**

All

**Total final enrolment**

309

**Key exclusion criteria**

Exclusion criteria based on self-report:

1. Bipolar disorder
2. Schizophrenia or other psychotic disorders
3. Personality disorders
4. Dementia and other neurodegenerative disorders
5. Heart surgery within the last 3 months
6. Untreated sleep apnea
7. Undergoing cancer treatment at the screening timepoint
8. Multiple sclerosis with an attack at screening timepoint
9. Adults without adequate skills in the Norwegian language to participate in the group-delivered course

**Date of first enrolment**

20/08/2022

**Date of final enrolment**

31/05/2023

**Locations****Countries of recruitment**

Norway

**Study participating centre****Alstahaug Healthy Life Centre**

Åsgata 13

Sandnessjøen

Norway

8800

**Study participating centre****Arendal Healthy Life Centre**

Østensbuveien 55

Arendal

Norway

4848

**Study participating centre**  
**Asker Healthy Life Centre**  
Nordre Bondi gård, Fredtunveien 25  
Asker  
Norway  
1386

**Study participating centre**  
**Bodø Healthy Life Centre**  
Speiderveien 4  
Bodø  
Norway  
8008

**Study participating centre**  
**Fredrikstad Healthy Life Centre**  
Jens Wilhelms gate 1  
Kråkerøy  
Norway  
1671

**Study participating centre**  
**Færder Healthy Life Centre**  
Øreveien 13  
Nøtterøy  
Norway  
3120

**Study participating centre**  
**Gausdal Øyer and Lillehammer Healthy Life Centre**  
Jørstadmovegen 690  
Fåberg  
Norway  
2625

**Study participating centre**  
**Jæren Healthy Life Centre**  
Ole Nielsensvei 20

Ålgård  
Norway  
4330

**Study participating centre**  
**Hamar Healthy Life Centre**  
Ankerskogvegen 7  
Hamar  
Norway  
2319

**Study participating centre**  
**Hole Healthy Life Centre**  
Furuhallveien 17B  
Helgelandsmoen Næringspark  
Røyse  
Norway  
3530

**Study participating centre**  
**Holmestrand Healthy Life Centre**  
Revåveien 18  
Sande  
Norway  
3070

**Study participating centre**  
**Inderøy Healthy Life Centre**  
Vennalivegen 7  
Inderøy  
Norway  
7670

**Study participating centre**  
**Kristiansand Healthy Life Centre**  
Kjøita Park 17 og 19  
Kristiansand  
Norway  
4630

**Study participating centre**  
**Lindesnes Healthy Life Centre**  
Lars O. Røllandsgate 7  
Mandal  
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4514

**Study participating centre**  
**Midt-Telemark Healthy Life Centre**  
Gullbring Kulturanlegg  
Gullbringvegen 34  
Bø  
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Furumoveien 70  
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**Ringerike Healthy Life Centre**  
Storgata 13  
Hønefoss  
Norway  
3510

**Study participating centre**  
**Skien Healthy Life Centre**  
Kongensgate 31  
Skien  
Norway  
3717

**Study participating centre**  
**Steinkjer Healthy Life Centre**  
Inn-Trøndelag helse- og beredskapshus, 5.et., Seilmakergata 10  
Steinkjer  
Norway  
7725

**Study participating centre**  
**Tromsø Healthy Life Centre**  
Seminarbakken 1  
Tromsø  
Norway  
9008

**Study participating centre**

**Trondheim Healthy Life Centre**  
Helse- og arenabygget Granåsen  
Smistadveien 13  
Trondheim  
Norway  
7026

**Study participating centre**  
**Verdal Healthy Life Centre**  
Ringveg Nord 1  
Verdal  
Norway  
7650

## Sponsor information

**Organisation**  
Norwegian University of Science and Technology

**ROR**  
<https://ror.org/05xg72x27>

## Funder(s)

**Funder type**  
Charity

**Funder Name**  
EkstraStiftelsen Helse og Rehabilitering (Stiftelsen Dam)

**Alternative Name(s)**  
Norwegian Foundation for Health and Rehabilitation, Dam Foundation, Stiftelsen Helse og Rehabilitering, EkstraStiftelsen Helse og Rehabilitering

**Funding Body Type**  
Private sector organisation

**Funding Body Subtype**  
Trusts, charities, foundations (both public and private)

**Location**  
Norway

# Results and Publications

## Individual participant data (IPD) sharing plan

The datasets generated during and/or analysed during the current study are not expected to be made publicly available due to participant confidentiality and considerations relating to data security. Participant-level data can be made accessible to specific users by contacting the project leader. The users must then be registered with the regional Medical Ethical Committee, in Norway. The need for restrictions may change over time, allowing the data to be made accessible at a later point.

## IPD sharing plan summary

Other

## Study outputs

| Output type                                   | Details                       | Date created | Date added | Peer reviewed? | Patient-facing? |
|---|-------------------------------|--------------|------------|----------------|-----------------|
| <a href="#">Results article</a>               |                               | 02/04/2025   | 14/04/2025 | Yes            | No              |
| <a href="#">Protocol article</a>              | protocol                      | 02/03/2023   | 03/03/2023 | Yes            | No              |
| <a href="#">Participant information sheet</a> |                               |              | 24/10/2022 | No             | Yes             |
| <a href="#">Participant information sheet</a> | Participant information sheet | 11/11/2025   | 11/11/2025 | No             | Yes             |
| <a href="#">Statistical Analysis Plan</a>     | version 1                     | 25/01/2024   | 19/02/2024 | No             | No              |
| <a href="#">Statistical Analysis Plan</a>     | version 2                     | 06/03/2024   | 08/03/2024 | No             | No              |
| <a href="#">Statistical Analysis Plan</a>     | version 3                     | 10/04/2024   | 15/04/2024 | No             | No              |
| <a href="#">Study website</a>                 | Study website                 | 11/11/2025   | 11/11/2025 | No             | Yes             |