A blended eHealth intervention for insomnia following acquired brain injury study protocol for a randomized controlled trial

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Abstract

Background Up to a third of stroke patients and patients with traumatic brain injury suffer from insomnia, including problems to fall asleep or stay asleep at night. Insomnia may exacerbate other brain damage-related problems, for example regarding cognitive functioning and emotional well-being, may lead to poorer quality of life, and may complicate recovery processes. Cognitive behavioural therapy for insomnia, delivered face-to-face or online, is found to be effective in the general population. However, despite the high prevalence and serious consequences of insomnia following acquired brain injury, studies on the efficacy of face-to-face cognitive behavioural treatment in this population are scarce, and this applies even more for studies on online cognitive behavioural therapy. Therefore, this study aims to evaluate the efficacy of a newly developed guided online cognitive behavioural therapy for insomnia following acquired brain injury. Methods A multicenter randomized controlled trial will be conducted, in which 48 patients diagnosed with stroke or traumatic brain injury, and insomnia will be randomly allocated to the online cognitive behavioural therapy for insomnia treatment group or the treatment as usual group. The treatment consists of 6 online cognitive behavioural therapy sessions given on a weekly basis and personalized feedback after each session, combined with face-to-face sessions. Outcomes will be assessed at baseline, immediately after the intervention period and at 6 week follow up. The primary outcome is the insomnia severity assessed with the insomnia severity index. Secondary outcome measures include sleep quality, sleep features derived from the sleep diary, fatigue, anxiety and depression, subjective cognitive functioning and societal participation. Discussion This study will provide insight on the efficacy of online cognitive behavioural therapy for insomnia following stroke and traumatic brain injury. Trial Register Netherlands Trial Register, NTR7082, 12 March 2018
Background

There is increasing awareness that sleep disorders following acquired brain injury are a serious problem and therefore need attention. These sleep disorders include sleep apnea, circadian rhythm disturbances, excessive daytime sleepiness, increased need for sleep, and insomnia (1-3). The latter is most commonly reported. Complaints include trouble falling asleep, staying asleep or waking up early and are accompanied by daytime complaints. Up to a third of stroke patients (4) and patients with traumatic brain injury (TBI) (2) meet DSM-IV criteria for insomnia disorder (ID), which is three times more than the 10% of the general population that suffers from ID (5). Insomnia may exacerbate other brain damage-related problems, for example regarding cognitive functioning and emotional well-being, may lead to poorer quality of life, and may complicate recovery processes (2, 4, 6). The treatment of insomnia should therefore be an important part of rehabilitation after acquired brain injury.

Numerous randomized controlled trials have shown that cognitive behaviour therapy for insomnia (CBT-I) is an effective treatment for insomnia in otherwise healthy people, as well as in several populations with insomnia comorbid with another disorder, (7-9). A problem with implementation of CBT-I on a large scale is the lack of skilled therapists, limiting access to treatment. Online versions of CBT-I have been developed to reach a larger group of patients with insomnia. Recent reviews show that internet delivered CBT-I is effective in improving sleep in adults with insomnia (10, 11). Despite the high prevalence and serious consequences of insomnia in stroke patients or patients with TBI, studies examining CBT-I following acquired brain injury are scarce. To date, CBT-I was examined in two studies concerning stroke patients (12, 13) and five studies concerning patients with mild to severe TBI (14-18). The CBT-I protocol applied in the studies consisted of four to eight weekly sessions, combining cognitive and
behavioural techniques including stimulus control, sleep restriction, cognitive therapy, and sleep hygiene education. CBT-I was adapted to brain injury patients by adding information on factors that could contribute to insomnia specifically after brain injury. CBT-I was administered online in one study (18) and face-to-face in the other six studies. Six out of these seven studies found significant improvement on sleep outcomes (12-14, 16-18) and four of these studies found significant improvement on secondary outcomes as depression and fatigue (12, 14, 15, 17). Only three studies were randomized controlled trials, of which two compared CBT-I with treatment as usual (12, 14) and one compared CBT-I with a placebo condition (online education only) (18). Furthermore, only three out of the seven studies – and none of the randomized controlled trials- formally assessed a diagnosis of ID as inclusion criterion (13, 16, 17). More randomized controlled trials using formal diagnosis of ID are needed in order to reach a well-founded conclusion on the efficacy of CBT-I in patients with stroke or TBI.

For the present study, we developed an online eHealth CBT-I (eCBT-I) (see box 1 for more details). Not only is eHealth easier to access and probably more cost-effective, it also has more benefits for patients, such as the opportunity to reread information and the freedom to follow treatment at their own time, place and pace. The present study will compare eCBT-I with treatment as usual in patients with stroke or TBI to evaluate the added value on top of standard care. Treatment as usual does not address sleep.

Objectives

The objectives of this study are:

(1) to evaluate the efficacy of eCBT-I in reducing insomnia severity post treatment and at six weeks follow up in patients with insomnia following a stroke or TBI, as compared to treatment as usual (TAU).

(2) to evaluate the efficacy of the eCBT-I in reducing complaints about cognitive
functioning, emotional well-being and societal participation posttreatment and at six weeks follow up compared to TAU.

(3) to evaluate whether treatment efficacies vary with severity of insomnia, with the severity and type of brain injury, and with the time since brain injury.

Methods

Study design

The study is a multicenter randomized controlled trial (RCT) comparing eCBT-I with TAU. A total of 48 patients will be randomly assigned to the eCBT-I or TAU group. Assessment will be performed at baseline (T1), one week after the 6-week intervention period (T2) and after 6-weeks follow up (T3). The total duration of participation is 14 weeks. See figure 1 for a flowchart.

Recruitment, randomisation, blinding and treatment allocation

Participants will be recruited from five outpatient rehabilitation centres spread over the Netherlands (Heliomare Rehabilitation at Wijk aan Zee, Reade at Amsterdam, Rehabilitation Friesland at Beetsterzwaag, Basalt Rehabilitation at Delft and The Hague and Adelante at Hoensbroek). Patients that are eligible based on the inclusion and exclusion criteria will be asked to participate. After informed consent is given, participants will complete the baseline measurement. Participants will then be randomly assigned to the eCBT-I or the TAU group. All participants will continue with standard rehabilitation care for various complaints, which do not specifically address insomnia. This care can include psychotherapy aimed at mood or other psychopathology, therapy aimed at cognitive functioning, physiotherapy, fitness, occupational therapy and social work. The TAU group can receive the eCBT-I after the study. Participants can leave the study at any time for any reason without consequences. In case of withdrawal, participants will be asked to still do the measurements to control for attrition bias and a new patient will be
included in accordance with the procedure. The measurements and intervention are not likely to have adverse consequences. Randomization will be done by a research randomizer program (www.randomizer.org) using blocks of four participants to balance participants equal to both groups within centres. No minimization procedure will be used. See table 1 for an overview of all measurements.

Eligibility criteria

Participants are eligible for inclusion if diagnosed with stroke or TBI and insomnia disorder according to DSM-5 criteria. Furthermore, they should be aged 18 or older and capable of using the internet. Exclusion criteria are untreated sleep apnea, current or expected treatment with a main focus on fatigue or sleep during the study, unstable medication regiments, use of medication with insomnia as side effect, alcohol or drug abuse, and a major untreated or unstable medical or psychiatric condition. Users of sleep medication will be encouraged to finish medication before enrolment or to keep intake stable during the study period.

Intervention

The eCBT-I comprises six guided weekly sessions provided online, combined with two face-to-face sessions of 60-minutes and a smartphone diary-app for daily reporting of sleep times and subjective sleep quality. Each session is structured around one topic and contains specific information, assignments and testimonials of two patients with insomnia after brain injury to illustrate sleep problems and homework assignments. All participants will receive online personal feedback after each session and will be encouraged to practice daily with the provided exercises, downloadable within the eHealth intervention on a daily basis. Participants can start with the next session after they have read their personal feedback. They can contact their therapist at any time by means of the integrated email function. Participants will be encouraged to complete the diary every day
by their therapist.

The eCBT-I is based on well-established CBT-I components and includes behavioural and cognitive techniques. These techniques contain sleep hygiene education, stimulus control, sleep restriction, cognitive restructuring, activation, relaxation, fatigue- and stress management. The eCBT-I has been adjusted to people with brain injury both with respect to content and the way of conveying information. Content adjustments include specific education about the nature and treatment of insomnia after brain injury, and adaptations to cognitive impairments due to brain damage. Information is given in clear and short texts and is visually supported. An option is included to allow for listening to an audio version of the texts. All sessions follow the same structure, with repetition of key points. Specific feedback suggestions for each session will be provided for the therapists.

The eCBT-I will be given by an experienced registered healthcare psychologist, trained in using the eHealth intervention. Adherence will be monitored by checking frequency of registration in the sleep diary app, time spent online, and online assignments done. For a detailed description of the intervention per week see Box 1.

Outcomes

Sleep outcome measures

The primary outcome measure is the change in insomnia severity measured with the Dutch version of the Insomnia Severity Index (ISI). The ISI consists of 7-items and uses a 5-point scale to measure to which extent participants experience insomnia. The total score ranges from 0 (no insomnia) to 28 (severe insomnia). A cut-off of 10 is used to indicate clinical levels of insomnia in this study, similar to other studies (19-21). The internal consistency is adequate (Cronbach’s alpha = 0.74-0.78). The ISI is selected as it is sensitive to treatment response (21, 22), and used in comparable research worldwide, including the Netherlands (19, 20). Secondary sleep outcome measures include overall
sleep disturbances assessed with the Pittsburgh Sleep Quality Index (23) and the following sleep features derived from the sleep diary app: total sleep time, sleep onset latency, number of nocturnal awakenings, sleep efficiency and subjective sleep quality.

Other outcome measures
These include fatigue after acquired brain injury assessed with the Dutch Multifactor Fatigue Scale (24); anxiety and depression assessed with the Dutch version of the 14-item Hospital Anxiety and Depression Scale (25); subjective cognitive functioning measured with the Cognitive Failure Questionnaire (26); and societal participation after rehabilitation assessed with the Utrecht Scale for Evaluation of Rehabilitation – Participation (USER-Participation) covering three aspects of societal participation: frequency of participation, restriction in participation and satisfaction with participation (27).

Other study parameters
Demographical, injury related and clinical variables which may influence the treatment effect will be registered: age, gender, diagnosis, time since injury, insomnia duration, use of prescribed sleep medication, use of other medication, comorbid psychiatric and somatic disorders, educational level, and currently being employed. Possible presence of sleep apnea will be screened with the Stop-Bang questionnaire (28) in participants that have not been evaluated for its presence with the gold standard overnight polysomnography (29).

Sample size calculation
In our latest randomized trial on the effect of eCBT-I we found an intraclass correlation coefficient (within subject correlation) of $r=0.54$ for ISI assessments repeated across 6 weeks in 175 people suffering from insomnia. Whereas treatment effects of eCBT-I are often reported to be of moderate to large size, we preferred to be somewhat conservative and expect a somewhat smaller than moderate effect in the current patient population.
Calculation of the required sample size using G*Power (30) indicated that 48 completers would provide, at a significance of alpha=0.05, sufficient power (1-beta=0.80) for a minimal detectable time-by-group interaction effect of $f=0.20$ (small to moderate).

**Statistical analyses**

Data analysis will be performed with SPSS 23 (IBM; Armonck, USA). Means and standard deviations of the demographic, injury related variables and the clinical characteristics collected at baseline will be calculated. Independent T-tests will be used to check for an imbalance between groups. Nominal variables will be checked with $\chi^2$-Tests and Ordinal variable with the Mann-Whitney Test. Statistical significance will be set at a p-value of 0.05. To accommodate likely occasional missing days, the repeated measures obtained by diaries will be analyzed with mixed effect models. The treatment effects will be examined using a repeated measures ANOVA. In the case of significant baseline imbalance, post hoc analysis of covariance (ANCOVA) will be used on the change scores of the outcome measures with baseline scores as covariates. Intention-to-treat analyses will be conducted. Additionally, a per protocol analyses will be performed. Also, we will compare the eCBT-I and TAU group with regard to the percentage of participants who have improved (reduction $> 7$ points on the ISI) and recovered (ISI$<10$). In an exploratory analysis, we will investigate whether the efficacy of the treatment varies as a function of severity of insomnia and severity, type of brain injury, and time since brain injury.

**Discussion**

This study is designed primarily to evaluate the effect of eCBT-I following stroke and traumatic brain injury, and therefore expected to provide insight on effects on sleep, fatigue, subjective cognitive functioning, emotional well-being and societal participation. It uses an online intervention that can be disseminated at low cost to professionals when found effective. Also, this study will explore who will benefit from treatment. As a result,
this study is expected to provide clinicians with evidence to help formulate future guidelines for the treatment of insomnia following stroke and TBI.

There are several limitations to this study that should be noted. In contrast to earlier research, we will combine stroke and TBI patients in our study, as insomnia complaints are comparable for those diagnoses. Our hypothesis is that eCBT-I will be similar effective for stroke and TBI patients. A potential limitation of this study is that differential effects between type of brain injury will be missed. This will be explored in post-hoc analysis. A second potential limitation is that we assess sleep only by subjective sleep measurements, as insomnia symptoms are our primary focus of interest. It would be interesting to use objective measures of sleep as is recommended as a standard research assessment of insomnia (31). However, we chose not to do this, as we expected this to be an additional burden on the participant that could negatively impact treatment adherence and increase drop-out.

Trial Status

Protocol version 2, September 19th, 2017. Recruitment started in January 2018 and is expected to be completed at December 2019. The final participants are expected to complete their assessments at the beginning of 2020.

Abbreviations

| Abbreviation | Description |
|--------------|-------------|
| ABI          | acquired brain injury |
| CBT-I        | cognitive behavioural therapy for insomnia |
| eCBT-I       | online cognitive behavioural therapy for insomnia |
| ID           | insomnia disorder |
| ISI          | insomnia severity index |
| TAU          | treatment as usual |
TBI traumatic brain injury

Declarations

Ethics approval

The study protocol, information brochure, questionnaires, and informed consent has received ethics approval from the Amsterdam University Medical Centre (AMC), protocol 2017-223. Central ethical approval at both central and local level has been confirmed from the Amsterdam University Medical Centre (ref approval no. 2017-223) and we will not begin recruiting at other centres in the trial until local ethical approval has been obtained. Informed consent will be obtained from all study participants. Study results will lead to public disclosure but can’t be traced back to the individual patients who took part in this study.

Consent for publication

Not applicable.

Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Disclosure statement

No potential conflict of interest is reported by the authors.

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Authors’ contributions

MF conceived the idea of this study and coordinates the study under supervision of EG, GG, CVB and EVS. EG, GG, and CVB contributed to the design of this study. MF and CVB
are involved in patient recruitment. MF was the primary author for this manuscript, and GG helped draft the manuscript. All authors critically reviewed the manuscript and approved the submitted version.

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Tables

Table 1 Assessment measures and time-points

| Assessment measure                                      | Enrolment | Baseline T=1 | Posttreatment T=2 | Follow-up |
|---------------------------------------------------------|-----------|--------------|-------------------|-----------|
| Eligibility screening                                   | X         |              |                   |           |
| Stop bang questionnaire                                 | X         |              |                   |           |
| Informed consent                                        | X         |              |                   |           |
| ISI                                                      | X         | X            |                   |           |
| PSQI                                                     | X         | X            |                   |           |
| Sleep diary                                             | X         |              |                   |           |
| DMFS                                                    | X         | X            |                   |           |
| HADS                                                    | X         |              |                   |           |
| CFQ                                                     | X         |              |                   |           |
| USER-P                                                   | X         |              |                   |           |
| Questionnaire received TAU                              |           |              |                   |           |
ISI: insomnia severity index; PSQI: Pittsburg sleep quality index, DMFS: Dutch multifactor fatigue scale; HADS: hospital anxiety and depression scale; CFQ: cognitive failure questionnaire; USER-P: Utrecht scale for evaluation of rehabilitation-participation; T=1: week 1; T=2: week 7; T=3: week 14

Box 1: Overview of online cognitive behavioural therapy for insomnia (eCBT-I)

| Week 1: | face to face session to provide information about the eHealth treatment and to optimize motivation for treatment |
|         | - start of online session 1: psychoeducation on sleep, the different sleep stages and sleep disorders acquired brain injury and their consequences in daily life. Homework assignment: map personal sleep prob their consequences for daily life together with coping so far: what was helpful and what was not? |
|         | - start with a daily sleep diary, which will be continued throughout the treatment. |
| Week 2: | online session 2: setting personal goals for treatment, information about sleep hygiene. Homework assign write down sub goals to improve sleep hygiene for the following week. |
| Week 3: | second face-to-face session to evaluate the personal goals for treatment |
|         | - online session 3: information on the relation between stress and sleep and different relaxation tech Homework assignment: practice of these relaxation techniques the following week. |
| Week 4: | online session 4: information on the circadian clock which is entrained by light and temperature and the inf activation on daytime sleepiness. Homework assignment: to balance activities and relaxation or to be more during daytime. |
| Week 5: | online session 5: different cognitive techniques, such as mindfulness and cognitive restructuring. Homework assignment: address and change unhelpful cognitive beliefs. |
| Week 6: | online session 6: consolidation and relapse prevention. |

Figures
Supplementary Files

This is a list of supplementary files associated with the primary manuscript. Click to download.

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