Nurse delivered Sleep Restriction Therapy for adults with Insomnia Disorder: process evaluation

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DOI: https://doi.org/10.3399/BJGP.2023.0162

To access the most recent version of this article, please click the DOI URL in the line above.

Received 03 April 2023
Revised 11 July 2023
Accepted 14 July 2023

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When citing this article please include the DOI provided above.
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Background

Sleep Restriction Therapy (SRT) is a behavioural therapy for insomnia.

Aim

To conduct a process evaluation of a randomised controlled trial comparing SRT delivered by primary care nurses plus sleep hygiene booklet with sleep hygiene booklet only for adults with Insomnia Disorder.

Design and setting

Mixed methods process evaluation.

Methods

We used semi-structured interviews of a purposive sample of patients receiving SRT, practice nurses delivering it, and general practitioners (GPs) or practice managers. Qualitative data were explored using Framework Analysis, and integrated with nurse comments and quantitative data, including baseline Insomnia Severity Index score and serial sleep efficiency outcomes to investigate relationships between these.

Results

We interviewed 16 patients, 13 nurses, 6 practice managers and 1 GP. Patients had no previous experience of behavioural therapy, needed flexible appointment times, and preferred face-to-face consultations; nurses felt prepared to deliver SRT, accommodating patient concerns, tailoring therapy, and negotiating sleep timings, despite treatment complexity and delays between training and intervention delivery. We explored how the intervention produced change, including patient and nurse interactions and patient responses to SRT. Difficulties maintaining SRT, negative attitudes towards treatment, and low self-efficacy were highlighted. Contextual factors, including freeing GP time, time constraints and conflicting priorities for nurses, with suggestions for alternative delivery options. Participants who found SRT a positive process showed improvements in sleep efficiency, whilst those that struggled did not.
Conclusion

SRT was successfully delivered by practice nurses and was generally well received by patients, despite some difficulties delivering and applying the intervention in practice.

How this fits in

Insomnia is the commonest sleep disorder and the second most prevalent mental health complaint in Europe with high economic costs due to health use, absenteeism and reduced productivity.

Lack of access to Cognitive Behavioural Therapy for Insomnia (CBT-I), which is recommended first line for insomnia, led to development and evaluation through a randomised controlled trial of Sleep Restriction Therapy (SRT), a component of CBT-I, delivered by primary care nurses.

This process evaluation of the HABIT trial of Sleep Restriction Therapy (SRT) with sleep hygiene compared with sleep hygiene alone, showed that SRT was successfully implemented with high fidelity by nurses and positively received by patients despite some initial difficulty adapting to changes in sleep schedules.

Introduction

Insomnia disorder, characterised by persistent problems with sleep initiation and/or maintenance and daytime consequences, is associated with mental and physical conditions and significantly impaired quality of life.[1] Affecting 10-12% adults, it is the commonest sleep disorder and the second most prevalent mental health complaint in Europe.[2]

Economic impacts of insomnia are high due to direct and indirect costs associated with increased healthcare usage, absenteeism, reduced productivity (‘presenteeism’) and accidents.[3]

Current guidelines recommend Cognitive Behavioural Therapy for Insomnia (CBT-I) as first line treatment,[4, 5] but lack of access to CBT-I in primary care [6] means GPs use sleep hygiene advice, hypnotics, or sedative antidepressants in preference.[7] Sleep restriction
therapy (SRT), a key ingredient of CBT-I, involves restricting and standardising a patient’s time in bed to address key perpetuating factors, including excessive or irregular time in bed, and daytime napping.[8] Meta-analysis suggests that SRT can reduce insomnia symptoms [9] but most trials have been performed in specialist research settings. In one small randomised control trial (RCT), a brief version of SRT significantly reduced insomnia severity compared to sleep hygiene alone at 6 months (Cohen’s d=0.54) when delivered in two sessions by a GP to a highly selected group of insomnia patients, free from comorbidities or medication use.[10] The need for a pragmatic trial testing a scalable model of treatment delivery in primary care, with a representative sample of people with insomnia, led to the design of the Health-professional Administered Brief Insomnia Therapy (HABIT) trial. HABIT was a parallel, open-label, RCT to determine the effectiveness of nurse-delivered SRT compared with sleep hygiene alone in adults (≥ 18 years) meeting criteria for Insomnia Disorder recruited within primary care.[11] Participants randomised to SRT received instructions from the nurse to keep a sleep diary, implement an agreed bed and rise-time, and calculate their sleep efficiency before each consultation. Patients received one consultation per week for four weeks, adjusting bed and rise times (sleep window) depending on reported improvements in sleep efficiency.[11] Nurse delivered SRT was found to clinically effective in reducing insomnia symptoms and improving sleep-related quality of life, depressive symptoms, mental health-related quality of life and work productivity at 3-, 6-, and 12-months follow-up.[12]. SRT was also highly likely to be cost-effective at a cost-effectiveness threshold of £20,000 per quality adjusted life year gained.[12]]

We conducted a process evaluation of the HABIT intervention in line with UK Medical Research Council (MRC) guidelines to understand intervention delivery, fidelity and acceptability from the perspective of patients, practice nurses and other stakeholders.[13] We aimed to explore how nurse-administered SRT in primary care worked, by examining implementation, mechanisms of impact, and contextual factors. ‘Implementation’ explores how the intervention is delivered and what is delivered, including the training and resources available as well as fidelity of delivery and any adaptations made. ‘Mechanisms of impact’ explore participants’ including perceived benefits and unintended or adverse effects. Exploring ‘contextual factors’ affecting implementation can help to understand the potential for sustaining and scaling the intervention more widely.
Methods

Design

We used a mixed methods design, integrating qualitative interview data and quantitative data collected from intervention participants. Data collection and analysis were conducted before the effectiveness and cost-effectiveness outcomes of the trial were revealed.

Qualitative interviews

Semi-structured interviews were undertaken with patients who had received SRT, nurses delivering the intervention and practice managers or GPs at participating practices using separate interview schedules (Boxes S1-3 respectively).

We aimed to interview 15 patients, 5 in each of the three trial areas, Oxford, Manchester and Lincolnshire. Patients were asked during their baseline assessment appointment for consent to be interviewed. Interviewees were selected from those consenting who had completed SRT within six months of interview, allowing us to explore how participants felt at various stages of the intervention. All nurses that delivered the intervention were contacted by the research team following consent and interviewed to explore their perceptions of providing SRT. Finally, practice managers or GPs from each participating practice were invited for interview, and those who consented were asked about their perceptions of impacts on the practice and the sustainability and scalability of the intervention. Interviews were conducted by telephone and were digitally recorded and transcribed.

Quantitative data

We compared patient interviewees’ perceptions of the intervention with two quantitative measures, baseline Insomnia Severity Index (ISI) and sleep efficiency recorded at each intervention session. ISI is a seven item self-reported questionnaire, scoring between 0-28, which assesses the severity, nature and impact of insomnia,[14] while sleep efficiency is the percentage of time in bed spent asleep (0-100%), which typically increases in participants for whom SRT is successful. Sleep efficiency was measured with a sleep diary over 7 days at baseline, and during the nurse intervention (recorded in nurse notes for treatment sessions
Fidelity assessments of audiotaped consultations were conducted by a clinical psychologist (DG) using a bespoke rating system and expressed as a percentage score.[11]

**Data analysis and integration**

Qualitative data were examined using Framework Analysis supported by NVivo 12. Two members of the team (SA and JP) conducted interviews and checked transcripts. Through familiarisation with the transcripts, examination of the interview schedules and the three MRC Framework domains, a set of *a priori* categories was developed to form an initial framework (Table 1).

Transcripts were coded independently (by SA and JP). An ‘other’ category was included in the framework to include relevant data that did not readily fit into the pre-existing categories. Whilst categories applied to each of the groups interviewed (nurse, patient and PM/GP), the codes were specific to each group as outlined in Table 1. Three researchers (SA, JP and NS), one of whom was independent of the initial analysis, discussed and agreed the final themes presented in the results.

Quantitative measures were presented at baseline (ISI and sleep efficiency) and at the nurse follow-up appointments (sleep efficiency) to show changes during treatment.

Relationships between qualitative findings, nurse records and quantitative measures were explored and presented using a joint display (Table S2) allowing us to directly compare the patient perceptions of SRT with changes to their measured sleep efficiency.

**Results**

We interviewed 16 patients, 13 nurses and 7 practice managers or GPs. Patients were aged from 19 to 74 (mean 56) years, including 7 male and 9 female interviewees, all of whom identified as White British. Patients are designated by region (A, B, C), sex (M, F) and age, e.g. Patient AF57. Nurses are designated by number and whether they were a CRN (Clinical Research Network) or practice nurse (Nurse).

Due to lack of availability of nurses at specific practices, two regions utilised research nurses (employed by their Local Clinical Research Network [LCRN]) rather than practice nurses. LCRN research nurses covered more than one practice and therefore 13 nurse participants
were interviewed. In two regions practices formed consortia under one management group, so 7 interviews were undertaken in the practice manager (six interviews) or GP (one GP interviewed) category.

Themes are listed under implementation, mechanisms of impact and contextual factors.

**Implementation of sleep restriction therapy**

The implementation category captured themes related to how the intervention was delivered, what was delivered and what the patients expected [13].

**Patients’ expectations due to lack of experience of behavioural therapy**

Patients did not know what to expect from SRT as most had no previous experience of behavioural therapy.

“No, it was the only sort of formal treatment I’ve had. I’ve tried things like relaxation, and things like that, but this was the only sort of scientific treatment I’ve had.” [Patient: AM57]

Patients hoped for improvements in their sleep pattern and daytime symptoms. They expressed how uncomfortable they felt if they had not slept well.

“I had the hope, rather than the expectation, that it would make me feel better in the morning; I would feel fresh, less tired.” [Patient: CF19]

**Feeling prepared, flexible appointment times, preferring face-to-face**

Nurses felt prepared and were supported with adequate training and tools enabling them to deliver SRT effectively.

“It was quite straightforward, and obviously we were provided with a PowerPoint presentation to go through; so that was really helpful” [Nurse 13]

Nurses and patients highlighted appointment flexibility as important. Face-to-face appointments were important for the initial appointment and maintaining motivation.

“So, I had a one-to-one meeting with a nurse, and I felt that those are really beneficial for me in terms of maintaining that treatment. For me personally, I don’t think I would have done it without the one to one.” [Patient: AM57]
“You absolutely can’t do the first one on the phone. [Although] From a patient perspective, it’s very convenient I guess, because they don’t have to come back to the practice.” [CRN Nurse 1]

**Accommodating concerns and tailoring therapy**

When patients had difficulty implementing SRT, particularly where their routines impacted on intervention delivery, nurses were able to modify SRT to the patient.

“There’s only one really out of the three, where I think there was a bit more tweaking of the times, if you like, and changing; purely because their routine was different”. [CRN Nurse-1]

“We tried to come up with a bit of a solution to it because not everybody is the same, so I felt it would be easier for me if I could knock it off in the morning. So, I didn’t mind getting up at 5.30 am rather than staying up.” [Patient: AF63]

SRT required individuals to calculate their sleep efficiency which was challenging for some patients. Nurses tailored sessions according to the patient’s ability to comprehend the process.

“It varied definitely. Some patients were able to engage very quickly, and the sessions could be done within 20 - 25 minutes because the patients were well engaged, able to understand the maths, able to understand what we wanted from them, how it was going to influence their sleep. Other patients, however, were very surprised about what they were expected to do, finding the concept very difficult.” [Nurse 5]

**Negotiating sleep timings**

Nurses negotiated bed and rise times with patients, as the protocol allowed for minor amendments to SRT to support a patient centred approach.

“And I said I was really struggling to get up at 5 am in the morning at the moment. So, we moved that to 5:15 last week.” [Patient: LNF51].

“I have actually played around with it (flexibility in sleep times) if they had been over 85%; particularly as I have got more used to it. I think initially when you start something; you get worried about how strict you have to be.” [Nurse 9]

**Learning to deliver despite complexity of SRT**
Initially, delivery and understanding of SRT involved a learning curve for both patients and nurses, who often adopted a collaborative approach to learning.

“It was a learning curve for both the nurse and myself; between us we worked out what was needed.” [Patient: CF65]

Nurses felt that the intervention was easy to deliver with practice.

“I think I was probably quite nervous to start with; but I think that is probably like most things, something new, and you do have teething problems when you start anything new”. [Nurse 2].

Patients suggested that they were able to calculate sleep efficiency but suggested that simplifying this might help retain people on the intervention.

“I expect other people would drop out because it took them a lot of time doing the calculations.” [Patient: AF55].

“It is very difficult to explain maths over the phone to a patient if they really struggle to understand it”. [Nurse 5]

**Challenge of delays**

For some nurses there were delays between training and seeing their first SRT patient, which increased the challenge of delivery.

“I think that was difficult because to do training and then wait, like quite a long time, till you are actually, physically seeing patients” [Nurse 4]

In one case nurses paired up to deliver SRT for their first patient to boost their. This minor divergence from the protocol was agreed in advance and might have been problematic if subsequent sessions were delivered by different nurses.

“I think myself and S doing it together, we seem to work quite well at this point, but as we get more patients, I think both of us will feel confident enough to do it on our own.” [Nurse 10]

**Mechanisms of impact of sleep restriction therapy**
We explored causal mechanisms, specifically how the delivered intervention produced change. We were interested in how participants interacted with nurses and responded to SRT and its effects.\[13\] This was crucial to understanding how the intervention worked.

**Self-motivation and effort**

Nurses observed that self-motivated patients were more likely to continue with SRT at home and those that put in the effort were more likely to succeed.

“The patients that have made it to the end of the study [end of intervention delivery] have taken it upon themselves to continue that process at home. I think it is because the patients that have made it through are self-motivated patients.” [CRN Nurse 1]

**Difficulty changing sleep habits**

Some patients tried hard to adhere with their SRT but changing their existing sleep habit was challenging.

“I had to stay up till midnight, and I thought – I’m never going to be able to do this. And I tried my hardest, but that was very difficult for me actually”. [Patient: AF63].

**Experiencing anticipated benefits**

Most patients reported that the initial week could be hard but, after that, they started to feel the benefits. They felt more refreshed, their sleep efficiency increased, and they were able to fall asleep more quickly and stay asleep.

“I mean after you get over that initial first week, you start to feel the benefits of it. I mean physically it hasn’t helped, because my condition, there’s not a cure for, but mentally I’m so much better for it, and it’s worth sticking with and seeing it through” [Patient: CF64].

Patients noted they fell asleep more quickly than prior to SRT and spent less time in bed awake. Nurses observed that patients receiving SRT perceived bedtime as a more positive experience and there were changes in the perception of sleep.

“Frequently could be anything up to an hour or an hour and a half previously, but now down to 15, 20, 25 minutes maximum, most nights before I drop off”. [Patient: CM65]
“She wasn’t having a nap, it was becoming a positive thing because she was looking forward to going to bed; and knowing that when she went to bed, she’d sleep. And even if she woke up, she said, she might wake up once or twice in the night, but she was able to get straight back off to sleep again. So that was good”. [CRN Nurse 1].

**Continuing support for adverse effects**

Patients did report some adverse effects during the initial phases of SRT.

“But the second week and third week, I felt exhausted. Really exhausted, so. Then felt alright again this week.” [Patient: CF51]

“I feel I could do with going to bed a bit earlier. I know in the booklet it suggests that you do things, but when you are so tired, you just can’t function.” [Patient: CF73].

Several patients and nurses highlighted the need for continued support following the end of the four-week therapy.

“Well for me anyway, four weeks wasn’t long enough for me at all…. What’s the point? So, I have reverted back to having naps now in the day. So, my insomnia at night has got worse.” [Patient: BF60].

“They were a bit like – ‘Where do we go from now?’...The chap I think was a little bit – ‘Oh!’; a little bit lost, if anything – ‘What do I do now?’; because he has not got anyone to report to at the end of the week. So just reassuring that he would get follow up at three months, six months.” [Nurse 3]

**Difficulties maintaining SRT**

Some patients expressed difficulties with very early rise times and the ability to maintain SRT every day.

“My hardest bit was the getting up at the time she wanted me to get up; and I couldn’t do that. I was getting up way too early, way too early.” [Patient: BF60].

**Reasons for withdrawal**

Nurses shared opinions of why patients were likely to withdraw from the intervention, which was related to conflicting commitments, tiredness, negative attitudes (in particular, where other commitments were perceived to be impacted), and lack of self-efficacy.
“Well, I can’t do it on a Saturday because of this, and I can’t do this or that.” [CRN Nurse 1].

“That was the biggest complaint that he just felt far too tired and didn’t feel he could go about his daily routines and things because of the tiredness.” [Nurse 2]

**Contextual factors in providing Sleep Restriction Therapy in primary care**

Practice managers, GPs and nurses all commented on contextual factors, relating to the practicalities of delivery within GP practices and the facilitators and challenges of sustaining and scaling up the intervention more widely.[13]

**Time constraints and conflicting priorities for nurses**

Practice managers were aware that nurses had concerns about time constraints. These included the difficulties of fitting in extended SRT appointments into existing consultation times which were generally shorter. There were also concerns about prebooking the appointments in advance, again due to lack of time.

“The nurse practitioners who are doing the study, they are enjoying doing it, but they are worried about time constraints; and in particular trying to get those four appointments booked in on a weekly basis. And in general practice, that’s very difficult for us” [Practice Manager 2]

**Freeing up GP time**

SRT could free up GP time, because it was an intervention that might stop patients calling into the surgery for sleep medication or to discuss their sleep problems.

“I think this is something that GPs would take on board quite readily because actually it’s taking work away from GPs and it’s giving an intervention that will actually free up time, I think, actually free up GP time. So if we can avoid patients phoning in for sleeping tablets, or coming to discuss sleep problems, and sort of following up these patients that goes on and on” [GP].

**Alternative delivery options**

Practice staff felt it would be helpful to designate specific times and days for the SRT clinic to be held. This would help staff organise clinics, book patients for appointments and free up time for nurses to complete additional administrative tasks associated with SRT delivery.
One suggestion was to consider treating SRT like other behaviour change clinics, including using set weekly times.

“The way I see it running is, if we treat it like a behaviour change intervention, just like our weight management courses” [Practice Manager 5]

Several practice managers wondered about using other staff members such as healthcare assistants.

“We have a very capable HCA [Health Care Assistant], who would be more than capable of actually sitting and going through this with someone; and obviously that would be a lot more cost effective”. [Practice Manager 2]

Small group therapy sessions were also suggested as a means of delivery and a way of optimising nurse time.

“I think it probably is good for the patients as well because as a group meeting for that education and going through it, there’s like a bit of a support group there for them as well.” [CRN Nurse 1].

“... for me I think a group environment with a nurse would have been just as effective as the one to one” [Patient: AF57].

“I think it probably be good for the patients as well because as a group meeting for that education and going through it, there’s like a bit of a support group there for them as well.” [CRN Nurse 1].

Practice staff, including GPs, were supportive, but had reservations about time constraints, availability, and set days for clinics. To ensure the intervention could be delivered in routine general practice suggestions were made that SRT is delivered in the format of other behavioural (e.g. smoking cessation and weight management) interventions.

Quantitative results and joint display

Most patients interviewed either had an improvement or at least no deterioration in sleep efficiency. Table S2 displays baseline ISI and baseline and subsequent sleep efficiency data together with summary extracts from any notes made by the nurses during the SRT sessions and a ‘representative’ quote from each patient regarding the SRT process. This indicated,
not unexpectedly, that participants who found SRT a positive process were more likely to show improvements in sleep efficiency, whilst those that struggled with SRT did not. Fidelity of nurse delivery was found to be high throughout the trial by the independent reviewer (audio recordings for session 1, median percentage fidelity score = 100%, and for session 3, median percentage fidelity score = 87.5%. [For full details see 12]
Discussion

Summary

In this process evaluation of a trial of Sleep Restriction Therapy (SRT) with sleep hygiene compared with sleep hygiene alone we found that SRT, despite its complexity, was successfully implemented with high fidelity by nurses and positively received by patients despite some initial difficulty adapting to changes in sleep schedules. We found that bed and rise times had to be negotiated and agreed with patients to enable these to be accepted and applied.

Strengths and limitations

This was a process evaluation of the largest pragmatic clinical trial of psychological therapy for insomnia to date delivered in primary care. A limitation of this study is that we interviewed participants that had mostly completed all 4 SRT sessions. We did speak to one participant that left the intervention midway due to an underlying health condition. A better understanding of why people withdrew from the intervention might inform changes to the intervention and ongoing support, leading to better retention.

Comparison with existing literature

Patients and nurses reported that they were able to quickly grasp the purpose of SRT and related processes. Patients preferred face-to-face consultations and felt that these helped maintain motivation. Although face-to-face interactions have been found to be preferred in some studies, overall the evidence is lacking that therapeutic alliance, disclosure, empathy, attentiveness or participation differs in face-to-face compared with telephone delivery of psychological interventions.[15] Some patients found calculating sleep efficiency difficult and felt that they needed help from the nurse, whilst nurses pointed out that helping someone with calculations over the telephone was harder than in person.

All patients interviewed found the first week of therapy difficult, due to reduced time in bed and strict bedtime and rising times. This is consistent with previous evaluations of SRT, where participants reported worsening of daytime functioning in the first week with improvements felt after a period of adjustment.[16, 17] It has been found that restriction of time in-bed time out-performs fixed bed and rise-times without restriction [18, 19]. This
suggests that whilst the initial increase in side-effects is challenging, they may also be a necessary part of the therapy linked to the need to harness homeostatic sleep pressure.[20, 21]

In this study there was negotiation between the nurses and the patients regarding sleep times and the need for flexibility, which was supported to some extent by the protocol.[11] Changing ingrained behaviours, in this case fixed night-time (or daytime nap) routines, was challenging and the flexibility on the part of the nurses allowed patients to feel some level of control. The flexibility was built into the protocol, and so did not affect the fidelity of delivery. One nurse interviewed did mention sharing delivery of the intervention with a colleague for one patient, which would only have been problematic if the patient was given inconsistent advice.

Participants reported adverse effects such as increased tiredness, ‘exhaustion’ and worries about driving which have been found in other studies.[17, 21] Others reported sleep disturbance due to menopause and use of sleep aids (e.g. sedatives) that affected allocated sleep time, which should be considered in future rollout of SRT.

Implications for practice and future research

Patients that did have improved sleep efficiency, also reported concerns, most commonly that four weeks of SRT was not long enough. All participants found the first week of the intervention very difficult as their body adjusted to limited time in bed. By the third week some were seeing significant benefits. For example, one participant who had improvements in sleep efficiency spoke of being woken by their alarm for the first time in years. Others only started to see benefits by the final week and as such felt the loss of support at the end of the intervention had a direct impact on their motivation to continue. Those that saw improvements earlier expressed being more likely to continue after treatment, while those that felt benefits later were more likely to revert to previous habits. One patient reported taking a nap in the afternoon the day after the final session and that they quickly reverted to their previous habits as there was no-one ‘watching over them’. This is a significant finding that indicated the importance of individual, personalised delivery with regular checks continuing for some until new habits and sleep patterns have been reinforced. It may
be relevant to think about cost effective refinements to SRT based upon these findings, such as extending SRT weekly sessions beyond 4 weeks.

Previous research has shown that it was possible for a GP to deliver an adapted version of SRT in general practice,[10] and this study confirmed that it was possible for practice nurses to consistently deliver the intervention. Practice managers and GPs also agreed that the intervention could be successfully delivered by nurses in this setting, which they considered may free up time for GPs. Suggestions to facilitate wider roll out included setting up specific clinics at set times that could be run by health care assistants and running small group sessions like other behaviour change (e.g. smoking cessation or weight loss) clinics, but this would require further evaluation.

Conclusions

SRT can be successfully delivered by nurses in general practice and was generally well received by patients. Ongoing support after the initial intervention period should be assessed to determine whether this leads to improved adherence and outcome.
Funding: The trial was funded by the National Institute for Health Research Health Technology Assessment Programme (HTA Project: 16/84/01).

Ethical Approval: Ethical approval was received from both Health Research Authority approval (IRAS: 238138) and ethical approval (Yorkshire and the Humber - Bradford Leeds Research Ethics Committee, reference: 18/YH/0153).

Competing Interests

Simon D. Kyle declares research funding from NIHR HTA (16/84/01 & 12/87/61), EME (131789) and the Oxford Health Biomedical Research Centre, and non-financial support from Big Health Ltd. in the form of no cost access to the digital sleep improvement programme, Sleepio, for use in clinical research (outside the submitted work). Paul Aveyard is NIHR Senior Investigator and declares research funding from NIHR HTA, NIHR Oxford Biomedical Research Centre, and NIHR Oxford and Thames Valley Applied Research Collaboration. Aloysius Niro Siriwardena declares research funding from Wellcome trust and NIHR HTA, RFPB and HS&DR. Ly-Mee Yu declares research funding from NIHR HTA. Peter Bower declares research funding from NIHR HTA. Leonie Maurer declares funding from NIHR Oxford BRC and consultancy fees from Mementor DE GmbH, outside the submitted work. Colin A. Espie declares research funding from NIHR HTA, EME and Oxford Biomedical Research Centre, and is co-founder of and shareholder in Big Health Ltd., a company which specialises in the digital delivery of cognitive behavioural therapy for sleep improvement (the Sleepio programme), outside the submitted work. Dimitri Gavriloff declares consultancy fees from Big Health Ltd., and Idorsia Pharmaceuticals Ltd., and is a previous salaried employee of Big Health Ltd. All other authors declare no competing interests.

Acknowledgements: We would like to acknowledge the participants who undertook the intervention, the nurses who delivered the intervention and the practice managers and GPs who supported the research within their practices.
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Table 1 Framework of categories and codes

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