A Multi-Component Cognitive-Behavioral Intervention for Sleep Disturbance in Veterans with PTSD: A Pilot Study

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Study Objectives: A significant portion of US military personnel are returning from deployment with trauma-related sleep disturbance, and disrupted sleep has been proposed as a mechanism for the development of medical conditions in those with posttraumatic stress disorder (PTSD). Although individuals with PTSD may realize improved sleep with either PTSD treatment or CBT for insomnia, many continue to experience residual sleep difficulties. Newly developed interventions designed to address nightmares are effective to this end, but often do not fully remove all aspects of PTSD-related sleep difficulties when used in isolation. A combined intervention involving both a nightmare-specific intervention and CBT for insomnia may lead to more marked reductions in PTSD-related sleep disturbances.

Methods: Twenty-two veterans meeting criteria for PTSD were enrolled in the study. A combined intervention comprised of CBT for insomnia and imagery rehearsal therapy was evaluated against a usual care comparison group.

Results: Intent-to-treat analyses revealed medium to large treatment effect sizes for all sleep diary outcomes, and very large treatment effects for insomnia severity, sleep quality, and PTSD symptoms.

Conclusions: Findings demonstrate that an intervention targeting trauma-specific sleep disturbance produces large short-term effects, including substantial reductions in PTSD symptoms and insomnia severity. Future research should focus on the optimal approach to the treatment of comorbid PTSD and sleep disturbance in terms of sequencing, and should assure that sleep-focused interventions are available and acceptable to our younger veterans, who were more likely to drop out of treatment.

Keywords: PTSD, insomnia, nightmares, imagery rehearsal therapy, nightmare rescripting, cognitive-behavioral therapy for insomnia, exposure, sleep quality


BRIEF SUMMARY

Current Knowledge/Study Rationale: CBT for Insomnia results in improved sleep across a variety of populations. Imagery Rehearsal Therapy (IRT) has been shown effective in those with post-traumatic stress disorder (PTSD). Addressing the sleep complaints of Veterans with PTSD may be optimized by combining CBT for Insomnia with IRT for nightmares.

Study Impact: Pilot study findings suggest that this combined intervention is promising for improving sleep and reducing PTSD symptoms in Veterans with PTSD. Given the increasing number of military personnel returning from deployment with PTSD and trauma-related sleep disturbance, trauma-specific sleep interventions are imminently needed.

Insomnia and nightmares are the two primary complaints of veterans with PTSD. Cognitive-behavioral therapy (CBT) for insomnia enjoys substantial empirical support for treating primary and comorbid insomnia.9 In veterans, a secondary analysis of data acquired from a larger treatment study showed that the subset with PTSD displayed pre-to-post treatment reductions in sleep onset latency and wake after sleep onset, and increased sleep efficiency and sleep quality in response to CBT for insomnia.10 PTSD patients showed a better overall subjective and objective response to CBT than they did to sleep hygiene education. In spite of these positive outcomes, PTSD
factors. While insomniacs look favorably upon sleep, those with trauma-related sleep disturbance are likely to view sleep as a necessary evil. Most veterans with PTSD experience hypervigilance; the need to remain aware of their surroundings at all times. Sleeping is an inherently contradictory experience to vigilance. Thus, veterans with PTSD are often sleep-avoidant, and this avoidance is further reinforced by the aversiveness of the nightmares that accompany sleep. Finally, the sleep architecture of individuals with PTSD differs from both normal sleepers and insomniacs, particularly with respect to the duration and onset of REM sleep. Taken together, these observations suggest that trauma-related sleep disturbance is an inherently different and more complex phenomenon than insomnia that is unlikely to remit using existing insomnia treatment approaches.

Several studies have presented data on behavioral treatments for nightmares in PTSD populations. Imagery rehearsal therapy (IRT), first presented in the empirical literature by Kellner and colleagues, was designed to reduce nightmare frequency and severity using “rescripting” of nightmares. Krakow and colleagues presented a more intensive conceptualization of this protocol in which nightmares are described as learned behaviors that can be relearned through the use of enhanced imagery skills. Imagery rehearsal therapy has been empirically validated and has produced favorable findings in terms of nightmares and PTSD symptoms in female sexual assault victims and deployed US army soldiers. IRT was combined with CBT for insomnia in two studies; a group-based intervention for crime victims with PTSD, and a single session behavioral protocol for adult victims of violent crime. Both produced improvements in nightmares, sleep quality and PTSD symptoms. Even more recently, Nappi and colleagues published a study where in IRT was employed as part of a VA clinical service. Treatment effect sizes (Cohen’s $d$) for nightmares, insomnia severity, and PTSD symptoms in the 35 veterans completing the course ranged from 0.16 (sleep quality) to 1.03 (PTSD symptoms).

Davis and colleagues developed a variation of IRT for nightmares that incorporates some standard treatment aspects of CBT-based PTSD treatment. Exposure, rescripting, and relaxation therapy (ERRT) was evaluated in a clinical trial of trauma-exposed adults (81.6% female, 67.3% meeting criteria for PTSD), and produced treatment effect sizes for nightmares, PTSD symptoms, self-reported sleep problems, and restfulness at one-week post-intervention ranged from 0.10 to 0.97 (Cohen’s $d$). Swanson and colleagues have since tested a combined version of ERRT and components of CBT for insomnia in 10 veterans with PTSD in a 10-session group therapy format. Treatment effect sizes for sleep and nightmares in this study ranged from 0.46 to 1.14 (Cohen’s $d$).

Given that CBT for insomnia and IRT have both been effective in reducing the sleep complaints of veterans with PTSD, our primary objective for the study was to assess the feasibility of a multi-component cognitive-behavioral sleep intervention for PTSD (SIP) using a randomized trial design. Our secondary purpose was to compare the intervention to usual care in terms of differential symptoms of sleep disturbance at post-intervention. We hypothesized that the combined effects of CBT for insomnia and IRT for nightmares would produce significantly greater improvements in sleep disturbance than usual care alone.

METHODS

Study Design

This study used a randomized parallel group experimental design. Participants were randomly assigned to either the intervention condition or a usual care condition, and were informed that the purpose of the study was to test the effectiveness of an intervention for improving sleep disturbance in veterans with PTSD. The study was approved by the institutional review board of our VA medical center, and all who enrolled provided written informed consent.

Participants

Study participants were recruited between 1/1/2008 and 12/31/2009 using flyers placed throughout the VA hospital and a community Veteran Center, and through letters sent to recently deployed veterans who had enrolled in a VA research registry and who had agreed to be recontacted for participation in VA research. To be considered for the study, veterans had to: (1) provide written informed consent; (2) meet DSM-IV-R criteria for a diagnosis of PTSD; (3) screen positive for an insomnia disorder on the Duke Structured Sleep Interview for Sleep Disorders; (4) score $> 14$ on the Insomnia Severity Index; and endorse nightmares on the PCL-M or the CAPs. Patients who screened positive on the DSISD for symptoms of sleep apnea, narcolepsy, restless legs syndrome, or circadian disorders, and those with active drug or alcohol abuse or dependence were excluded. All participants were required to have a PTSD diagnosis established using the Clinician-Administered PTSD Scale (CAPS) or the SCID. Of the 93 self-referred prospective study participants, 15 men and 7 women met study selection criteria, completed baseline procedures, and were subsequently randomized to treatment conditions (Figure 1). Of the 22 who were ultimately randomized to condition, PTSD diagnosis was established in 20 using the CAPs and in 2 using the SCID.

The average age of participants was 45.96 (SD 11.06). Most participants (73%) were either African American (N = 8) or Caucasian (N = 8). The demographic and mental health characteristics of participants are outlined in Table 1. Enrolled study participants were allowed to be engaged in mental health treatment during the study period. Table 2 outlines the specific type and frequency of mental health contacts received by each study participant during the study period.

Measures

Screening and Outcome Measures

Duke Structured Interview for Sleep Disorders (DSISD): The DSISD is an instrument developed to assist in ascertaining DSM-IV and International Classification of Sleep

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Disorders sleep disorder diagnoses. The DSISD includes questions that incorporate criteria for ascertaining sleep disorders within both the DSM-IV and the recently updated ICSD sleep disorder nosologies. The instrument has acceptable reliability and discriminant validity, and is effective for discerning the types of sleep disorders that would disqualify participants for this study.

Folstein Mini-Mental Status Exam: The Folstein Mini-Mental Status Exam (MMSE) was administered to all study candidates using standard administration/scoring procedures. The MMSE was used to identify and exclude individuals who have cognitive deficits that would preclude their ability to provide informed consent or to fully participate in an interactive treatment process. As consistent with clinical applications for dementia screening, those with a score < 24 were excluded from the study.

Insomnia Severity Index (ISI): The ISI is a 7-item questionnaire that provides a global measure of perceived insomnia severity. Each item is rated on a 5-point Likert scale, and the total score ranges from 0-28. The following guidelines are recommended for interpreting the total score: 0-7 (no clinical insomnia), 8-14 (subthreshold insomnia), 15-21 (insomnia of moderate severity), and 22-28 (severe insomnia). The ISI has adequate psychometric properties, has been validated against diary and polysomnographic measures of sleep, and has been shown sensitive to therapeutic changes in several of our treatment studies of insomnia. The ISI was used to determine treatment eligibility, to assess treatment outcome, and to determine clinical significance of study findings.

Electronic Sleep Diary: Subjective sleep estimates were obtained using a hand-held computer (PDA) containing an interactive program that automates the collection of subjective sleep data. The PDA device was programmed to elicit daily responses from participants and electronically record multiple days of subjective sleep information, in addition to the number and severity of nightmares for the previous night. Five variables of interest were calculated or obtained from electronic sleep diaries, as follows: total sleep time (TST); sleep onset latency (SOL); wake after sleep onset (WASO); sleep efficiency % (SE); and nightmare frequency (NM frequency).

The Pittsburgh Sleep Quality Index (PSQI): The PSQI is a self-rating scale that yields a quantitative index of general sleep quality/disturbances. The PSQI is composed of 4 open-ended questions and 19 self-rated items (0-3 scale) assessing sleep quality and disturbances over a 1-month interval, and yields a
a sensitivity of 86% and specificity of 78% for detecting major depression, with a score ≥ 2 correctly categorizing the greatest number of those with major depression.

Procedure

Patients expressing interest in the study were first screened for study eligibility by phone, with the exception of a few veterans who presented to the office of the PI in response to flyers. A more rigorous study screening assessment was conducted in person. Those meeting study eligibility criteria were enrolled in the study, completed a baseline demographics questionnaire, and then conducted one week of sleep monitoring at home using the electronic sleep diary. Upon returning with study equipment, they completed a baseline packet of questionnaires assessing study outcome variables. They were then randomized to condition and to therapist. Those in the intervention condition were scheduled for their first session of SIP within 3 weeks and completed 6 bi-weekly intervention sessions over the following 12 weeks. Following their sixth session of SIP, they were again sent home with equipment to monitor sleep and then completed the post-intervention questionnaire packet upon their return. The usual care control condition procedures were identical, excepting that they did not receive the intervention. All usual care participants were offered the opportunity to receive SIP at the end of the study. All participants were paid $40 for each of the 2 assessment periods and $15 for study screening.

Therapists and Treatments

Treatment was conducted by the PI (CSU) and the co-investigator (JDE), who are both licensed clinical psychologists. Of the 13 randomized to the intervention condition, CSU treated 5 and JDE treated 8.

Usual Care/Usual Care Control: Veterans seeking assistance for sleep disturbance and PTSD symptoms at the VA are often treated for their symptoms by their primary care provider (PCP). PCPs may provide hypnotics, antidepressants, anxiolytics, and mood stabilizing medications to address veteran symptoms, in addition to referring the veteran to the mental health clinic.

Sleep Intervention for PTSD (SIP): Patients in the SIP condition were eligible to receive the same elements as the Usual Care patients. In addition, these patients received 6 bi-weekly 1-h individual sessions with the interventionist, including 3 sessions of cognitive-behavioral therapy (CBT) for insomnia and 3 sessions of imagery rehearsal therapy (IRT), in that order. CBT for insomnia consisted of a prescription for an individually tailored behavioral regimen based upon sleep restriction theory, stimulus control, standard sleep hygiene recommendations, and the identification and restructuring of dysfunctional beliefs and attitudes regarding sleep. The IRT component included education about the role of learning in nightmares, visual imagery skill building, and specific instructions on how to rescript nightmares. Participants were instructed to change the nightmare in any way they liked and to practice this technique for ≥ 15 min/day, and to practice rescripting no more than 2 new dreams per week.

Analyses

Baseline group differences were assessed using standard χ² procedures for categorical measures, and the F distribution for continuous variables.
Intent-to-Treat Outcome Analyses

For all intent-to-treat outcomes, linear mixed-effects models (PROC MIXED in SAS) were used to determine expected mean values at each time point and to test hypotheses of group differences. The model included time and the group-by-time interaction; an unstructured covariance matrix was used to account for the within-patient correlation over time. All available data, including data from participants who subsequently discontinued the study, were used for the longitudinal analyses. Mixed-effects models assume non-informative dropout, meaning that the probability of dropout may depend on covariates or a participants’ previous responses but not on current or future responses. A p value < 0.05 was considered statistically significant. Due to a PDA programming error, we lost a significant portion of the data for nightmare severity, so this variable was not included in analyses.

Completer Outcome Analyses

To analyze outcomes for completers only, a series of ANOVAs (analysis of variances) involving one between-subjects factor (SIP + Usual Care vs. Usual Care Only) and one within-subjects factor (time) were used to assess for group by time interactions. Since one study participant in the intervention group did not provide post-intervention sleep log data, these analyses were conducted on 9 usual care participants and 8 intervention participants. A p value < 0.05 was considered statistically significant.

Effect Size Calculations

Effect sizes were calculated for each outcome by subtracting post-intervention values for the usual care group from post-intervention values for the treatment group, and then dividing by the pooled baseline standard deviation.

Test of Clinical Significance

To assess the clinical significance of our findings, we employed the criteria used by Morin and colleagues, with remission from insomnia defined as a post-intervention ISI score ≤ 7 (no insomnia), and response defined as a 8-point drop in ISI score (a categorical change) from pre- to post-assessment. As discussed above, remission from PTSD was defined as a PCL-M score ≤ 49; sleep quality remission was defined as a post-intervention PSQI score ≤ 5; and remission from PTSD-specific disruptive nocturnal behaviors was defined as a PSQI-A post-intervention score < 4. There are no established criteria for sleep diary values reflecting “normal” sleep in those with PTSD. However, a criterion of < 31 min of WASO or SOL was found to discriminate those with insomnia from normal sleepers. Since individuals with PTSD endorse high levels of insomnia, we utilized this criterion as being reflective of normal WASO and SOL at post-intervention. Treatment response status was evaluated using χ² analyses, wherein the frequency of responders and remitters, as defined above, were compared across conditions.

RESULTS

Treatment Completers versus Drop-Outs

Four of the 22 randomized study participants dropped out of the intervention group prior to completing the study. There were no dropouts from the usual care group. The 4 participants who did not complete the study were all male OEF/OIF veterans. They were also younger (F₁,20 = 7.47, p = 0.01, Drop Out M = 34, SD = 2.94, Completer M = 48.61, SD = 10.42), had greater sleep onset latency (F₁,20 = 5.76, p = 0.03, Drop Out M = 83.93, SD = 52.66, Completer M = 42.95, SD = 25.15), reported less restful sleep on diaries (F₁,20 = 5.51, p = 0.03, Drop Out M = 2.5, SD = 0.37, Completer M = 4.08, SD = 1.31), and had poorer sleep quality (higher PSQI scores) (F₁,19 = 4.69, p = 0.04, poorer sleep quality) than study completers at baseline. Two veterans dropped out due to work conflicts, one because of the distance from his residence to the VA, and one was lost to follow-up. Those not completing the study did not otherwise differ from completers on demographic characteristics or outcome measures.

Baseline Comparisons

Groups were compared on baseline demographic and outcome characteristics using univariate ANOVA analyses and χ² analyses, as appropriate, for those completing the study. Groups did not differ on age, ethnicity, or gender distribution, time since trauma, or the number of Criterion A traumatic events (Table 1). In addition, participants did not differ across groups on any baseline outcome variables (Table 3). Three variables were considered as possible covariates: age, time since trauma, and baseline anxiolytic/hypnotic medication use. Since none of these variables correlated with baseline outcome measures, they were not included as covariates in analyses.

Therapist Effects

A series of repeated measures ANOVAs were conducted to assess for therapist effects for each outcome measure (Outcome Measure × Group × Therapist). No therapist effects or group by therapist effects were found for any outcome measures.

Mental Health Treatment Group Effects

Chi-square analyses were used to assess for group differences on treatment engagement during the study protocol. Groups did not differ on the percentage engaged in mental health treatment (χ² = 0.90, p = 0.34) or medication management (χ² = 2.10, p = 0.15) during the study period (See Table 2).

Table 3—Baseline sleep characteristics

<table>
<thead>
<tr>
<th>Measure</th>
<th>SIP</th>
<th>Usual Care</th>
<th>Difference Statistic</th>
</tr>
</thead>
<tbody>
<tr>
<td>TST</td>
<td>315.45</td>
<td>311.99</td>
<td>0.01</td>
</tr>
<tr>
<td>WASO</td>
<td>85.07</td>
<td>103.83</td>
<td>0.71</td>
</tr>
<tr>
<td>SOL</td>
<td>50.30</td>
<td>50.54</td>
<td>0.00</td>
</tr>
<tr>
<td>SE</td>
<td>70.52</td>
<td>66.47</td>
<td>0.54</td>
</tr>
<tr>
<td>NM</td>
<td>0.72</td>
<td>0.59</td>
<td>0.32</td>
</tr>
<tr>
<td>Frequency</td>
<td>22.77</td>
<td>22.00</td>
<td>0.17</td>
</tr>
<tr>
<td>ISI</td>
<td>63.04</td>
<td>64.44</td>
<td>0.01</td>
</tr>
<tr>
<td>PCL-M</td>
<td>3.77</td>
<td>3.89</td>
<td>0.02</td>
</tr>
<tr>
<td>PHQ</td>
<td>14.17</td>
<td>14.33</td>
<td>0.01</td>
</tr>
<tr>
<td>PSQI</td>
<td>10.00</td>
<td>10.00</td>
<td>0.00</td>
</tr>
</tbody>
</table>

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Questionnaire Outcomes

Intent-to-treat statistical analyses showed a significant group by time interaction in favor of SIP over the usual care condition for most questionnaire outcomes. SIP produced significantly greater improvements in insomnia severity (ISI) ($F_{1,21} = 11.80, p = 0.003$), PTSD symptoms (PCL-M) ($F_{1,21} = 22.72, p = 0.0001$), and sleep quality (PSQI) ($F_{1,21} = 17.31, p = 0.0005$) (Table 6). Groups did not differ at post-intervention on depression (PHQ-2) or the PTSD-specific sleep quality measure (PSQI-A). The effect size for all significant questionnaire outcomes fell in the "large" range, and in the "small to moderate" range for nonsignificant questionnaire outcomes.

ANOVA As were then conducted to assess for group differences for those with complete baseline and post-intervention data on the amount of change in their scores on self-report questionnaires (ISI, PCL-M, PHQ, PSQI, and PSQI-A) from baseline to post-intervention (Table 7). ANOVAs revealed a significant time by group interaction for the ISI ($F_{3,14} = 9.88, p = 0.006$), the PCL-M ($F_{3,14} = 17.63, p = 0.001$), and the PSQI ($F_{3,14} = 15.91, p = 0.001$), with those in the intervention condition faring better at post-intervention than the usual care condi-

### Table 4—Comparison of predicted means, standard error (SE) values and treatment effect sizes for intent-to-treat groups on sleep diary outcomes

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline</th>
<th>Post-Intervention Predicted</th>
<th>Difference Statistic</th>
<th>Effect Size</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>SIP Group</td>
<td>TAU Group</td>
<td>F</td>
</tr>
<tr>
<td>TST (Hours)</td>
<td></td>
<td>5.23</td>
<td>6.47</td>
<td>5.26</td>
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<tr>
<td></td>
<td></td>
<td>0.24</td>
<td>0.37</td>
<td>0.35</td>
</tr>
<tr>
<td>WASO (Minutes)</td>
<td></td>
<td>92.75</td>
<td>42.03</td>
<td>88.00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10.87</td>
<td>14.47</td>
<td>13.73</td>
</tr>
<tr>
<td>SOL (Minutes)</td>
<td></td>
<td>50.40</td>
<td>22.39</td>
<td>45.03</td>
</tr>
<tr>
<td></td>
<td></td>
<td>7.30</td>
<td>6.55</td>
<td>6.32</td>
</tr>
<tr>
<td>SE (Percentage)</td>
<td></td>
<td>68.86</td>
<td>86.89</td>
<td>70.91</td>
</tr>
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<td></td>
<td></td>
<td>2.68</td>
<td>3.24</td>
<td>3.07</td>
</tr>
<tr>
<td>NM Frequency</td>
<td></td>
<td>0.67</td>
<td>0.51</td>
<td>0.83</td>
</tr>
<tr>
<td>(Mean per Night)</td>
<td></td>
<td>0.11</td>
<td>0.18</td>
<td>0.17</td>
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</tbody>
</table>

*p < 0.05, **p < 0.01

### Table 5—Comparison of means, standard error (SE) values and treatment effect sizes for participants with complete baseline and post-intervention data on sleep log outcomes

<table>
<thead>
<tr>
<th>Sleep Log Variable</th>
<th>Baseline</th>
<th>Post-Intervention</th>
<th>ANOVA Statistics</th>
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<tr>
<td></td>
<td></td>
<td>SIP N = 8</td>
<td>TAU N = 9</td>
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<tr>
<td>TST (Hours)</td>
<td></td>
<td>5.27</td>
<td>6.56</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.29</td>
<td>0.46</td>
</tr>
<tr>
<td>WASO (Minutes)</td>
<td></td>
<td>88.93</td>
<td>30.95</td>
</tr>
<tr>
<td></td>
<td></td>
<td>10.58</td>
<td>7.34</td>
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<tr>
<td>SOL (Minutes)</td>
<td></td>
<td>42.95</td>
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<td></td>
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<td>5.93</td>
<td>2.11</td>
</tr>
<tr>
<td>SE (Percentage)</td>
<td></td>
<td>69.98</td>
<td>89.58</td>
</tr>
<tr>
<td></td>
<td></td>
<td>3.12</td>
<td>2.06</td>
</tr>
<tr>
<td>NM Frequency (Mean per Night)</td>
<td></td>
<td>0.63</td>
<td>0.27</td>
</tr>
<tr>
<td></td>
<td></td>
<td>0.53</td>
<td>0.26</td>
</tr>
</tbody>
</table>

*p < 0.05, **p < 0.01

### Sleep Diary Outcomes

Intent-to-treat statistical analyses showed a significant group by time interaction in favor of SIP over the usual care condition for all sleep diary outcomes (Table 4). SIP produced significantly greater baseline to post-intervention improvements in sleep diary measures of TST ($F_{1,21} = 6.33, p = 0.02$), WASO ($F_{1,21} = 5.91, p = 0.02$), SOL ($F_{1,21} = 10.17, p = 0.004$), SE ($F_{1,21} = 14.61, p = 0.001$), and nightmare frequency ($F_{1,21} = 5.03, p = 0.03$, $p = 0.04$) (Table 4). The effect sizes for all sleep diary outcomes fell in the “medium” to “large” range using Cohen’s original conceptualization of these terms.13

Completer analyses revealed a significant group by time interaction for nightmare frequency ($F_{1,15} = 4.96, p = 0.04$), with those in the intervention group reporting a significantly greater reduction in nightmares from baseline to post-intervention than the usual care group (Table 5). Trends toward significance were found for total sleep time ($F_{1,15} = 3.54, p = 0.08$) and sleep efficiency percentage ($F_{1,15} = 3.70, p = 0.07$), with the intervention group reporting more total sleep time and a higher sleep efficiency at post-intervention relative to usual care. Effect sizes for all completers-only sleep log outcomes fell in the “large” range.

### Questionnaire Outcomes

ANOVA As were then conducted to assess for group differences for those with complete baseline and post-intervention data on the amount of change in their scores on self-report questionnaires (ISI, PCL-M, PHQ, PSQI, and PSQI-A) from baseline to post-intervention (Table 7). ANOVAs revealed a significant time by group interaction for the ISI ($F_{3,14} = 9.88, p = 0.006$), the PCL-M ($F_{3,14} = 17.63, p = 0.001$), and the PSQI ($F_{3,14} = 15.91, p = 0.001$), with those in the intervention condition faring better at post-intervention than the usual care condi-
The intervention group participants (4/4 or 100%) in this subsample had achieved “normal” SOL as compared to only 14% of usual care participants (1/7) ($\chi^2 = 7.5$, $p = 0.02$). Groups did not differ at post-intervention on the percentage of WASO remitters ($\chi^2 = 0.79$, $p = 0.55$), with 1 of 9 remitting in the wait-list group and 2 of 7 in the intervention group.

### Clinical Significance Findings

#### Sleep Diary Outcomes

Post-assessment sleep diary data was not available for 1 participant in the intervention group. Figures 2 and 3 depict baseline to post-assessment change in SOL and WASO in those providing complete sleep diary data. As depicted, 4 subjects in the intervention group had baseline SOL values < 31 min, and one subject had a baseline WASO value < 31 min. In the waitlist group, 2 participants had SOL values < 31 min. Baseline SOL and WASO values did not differ between groups when considering only those participants with complete sleep diary data and baseline SOL and WASO values above the criterion level for “normal” sleep. At post-intervention, however, all of the intervention group participants (4/4 or 100%) had achieved “normal” SOL compared to only 14% of usual care participants (1/7) ($\chi^2 = 7.5$, $p = 0.02$). Groups did not differ at post-intervention on the percentage of WASO remitters ($\chi^2 = 0.79$, $p = 0.55$), with 1 of 9 remitting in the waitlist group and 2 of 7 in the intervention group.

### Questionnaire Outcomes

Figures 4, 5, and 6 depict baseline to post-intervention change in ISI, PCL-M, and PSQI scores, respectively. At post-intervention, one participant in the intervention group remitted from insomnia (1/9 or 11%), and 4 were insomnia responders (4/9 or 44.4%). As depicted in Figure 5, 1 participant in each group had a baseline PCL-M score < 50. Baseline PCL-M values did not differ between groups when considering only those participants with complete sleep diary data and baseline PCL-M values above the PTSD screening cutoff of 50 ($F = 0.17$, $p = 0.69$). At post-intervention, however, half of the intervention group participants in this subsample (4/8 or 50%) had remitted from PTSD as compared to none of the usual care participants (0/8) ($\chi^2 = 5.33$, $p = 0.04$). In terms of

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Table 6—Comparison of predicted means, standard error (SE) values, and treatment effect sizes for intent-to-treat groups on questionnaire outcomes

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline</th>
<th>Post-Intervention Predicted</th>
<th>Difference Statistic</th>
<th>Effect Size</th>
</tr>
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<tbody>
<tr>
<td></td>
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<td>TAU Group</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>M</td>
<td>SE</td>
<td>F</td>
</tr>
<tr>
<td>ISI</td>
<td></td>
<td>22.46</td>
<td>12.45</td>
<td>21.58</td>
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<tr>
<td>PCL-M</td>
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<td>62.39</td>
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<td>66.08</td>
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<td>PHQ</td>
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<td>3.81</td>
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<td>4.06</td>
</tr>
<tr>
<td>PSQI</td>
<td></td>
<td>14.24</td>
<td>9.31</td>
<td>14.47</td>
</tr>
<tr>
<td>PSQI-A</td>
<td></td>
<td>10.00</td>
<td>7.74</td>
<td>9.11</td>
</tr>
</tbody>
</table>

*p < 0.05, **p < 0.01

---

Table 7—Comparison of means, standard error (SE) values and treatment effect sizes for participants with complete baseline and post-intervention data on outcome questionnaires

<table>
<thead>
<tr>
<th>Measure</th>
<th>Baseline</th>
<th>Post-Intervention</th>
<th>ANOVA Statistics</th>
<th>Group (A)</th>
<th>Time (B)</th>
<th>A × B</th>
<th>Cohen's d</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>SIP N = 9</td>
<td>TAU N = 9</td>
<td>F</td>
<td>F</td>
<td>F</td>
<td></td>
</tr>
<tr>
<td>ISI</td>
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<td>22.22</td>
<td>12.44</td>
<td>21.44</td>
<td>11.80**</td>
<td>-2.15</td>
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<td>PCL-M</td>
<td></td>
<td>62.39</td>
<td>45.25</td>
<td>66.08</td>
<td>22.72**</td>
<td>-1.76</td>
<td></td>
</tr>
<tr>
<td>PHQ</td>
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<td>3.45</td>
<td>4.06</td>
<td>0.86</td>
<td>-0.34</td>
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</tr>
<tr>
<td>PSQI</td>
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<td>14.24</td>
<td>9.31</td>
<td>14.47</td>
<td>17.31**</td>
<td>-1.60</td>
<td></td>
</tr>
<tr>
<td>PSQI-A</td>
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<td>10.00</td>
<td>7.74</td>
<td>9.11</td>
<td>0.76</td>
<td>-0.30</td>
<td></td>
</tr>
</tbody>
</table>

*p < 0.05, **p < 0.01
Figure 2—Sleep onset latency

Figure 3—Wake after sleep onset

Figure 4—Insomnia severity
Sleep quality, groups did not differ at post-intervention on the percentage of sleep quality remitters, with 3 (3/9 or 33.3%) of those in the intervention group remitting, and none of those in the usual care group (0/9 or 0%). Finally, with regard to PTSD-specific disruptive nocturnal behaviors, groups did not differ at post-intervention with 3 (3/9 or 33.3%) of those in the intervention group remitting and one (1/9 or 11.1%) in the usual care group ($\chi^2 = 1.29$, $p = 0.26$).

**DISCUSSION**

The primary purpose of our study was to assess the feasibility of SIP using a randomized trial design. Although the study revealed the need for slight procedural changes to be incorporated into a larger trial, the overall findings suggest that the intervention is feasible and generally acceptable to veterans with PTSD. Our secondary purpose was to compare the intervention to usual care in terms of differential symptoms of sleep disturbance at post-intervention. We hypothesized that this combined intervention, in addition to usual care, would produce better outcomes than usual care alone. Our hypothesis was supported in terms of both average improvement across individual measures and the clinical significance of our findings. Intent-to-treat analyses revealed medium to large effect sizes (group differences) for all sleep diary outcomes, and very large treatment effects (Cohen’s $d > 1.50$) for insomnia severity, sleep quality, and PTSD symptoms. In contrast, none of the participants in the usual care group responded or remitted from insomnia or PTSD, and did not improve from baseline on sleep quality. SIP did not produce a treatment effect for depression, and although SIP is not designed to treat depression, this finding was surprising in light of the treatment effect on correlated issues: insomnia and PTSD. We suspect that the failure to find a significant effect for depression may be related to the restricted
In spite of large effect sizes, remission rates were lower than hoped for measures of both insomnia (11%) and sleep quality (33%). In fact, it is particularly surprising that remission rates for sleep targets were lower than for PTSD (50%). Comparison of PSQI and ISI items across groups does not reveal a consistent indicator of residual symptomatology that might explain the low remission rates for insomnia and sleep quality. However, of all sleep log variables assessed, the treatment effect sizes were lowest for nightmare frequency. Since there was no follow-up assessment included in the design of this pilot study, it remains unknown if the IRT skills acquired during the intervention might have resulted in better outcomes at a later follow-up assessment, or if decrements in clinical improvements would be revealed over time. However, it is plausible that residual nightmares in the treatment group might explain the low remission rates for insomnia and sleep quality. This explanation is also consistent with our finding of greater remission rates for SOL than WASO on sleep logs.

Randomized controlled trials of sleep-focused interventions in veterans with PTSD are largely absent from the empirical literature. However, our findings are generally consistent with the available studies addressing post-trauma sleep disturbance, excepting the larger effect size produced by SIP relative to other interventions. SIP produced significant treatment effects in the same domains as those found in the clinical outcomes of IRT for veterans described by Nappi and colleagues, including nightmares, insomnia severity, and PTSD symptoms, with the exception that we also found a significant treatment effect for sleep quality. Our findings are also generally consistent with those of Davis et al. ERRT resulted in a significant improvement in depression, however, whereas depression did not improve significantly with SIP, as discussed above. Insomnia was not assessed, and sleep diaries were not included in the study by Davis et al., so no comparisons can be made here.

SIP differs from previously assessed interventions in at least two areas. First, exposure is not an intended mechanism of change in SIP, in contrast with empirically based PTSD treatments and ERRT. In spite of differences in treatment protocols, however, SIP (50%) and ERRT (46%) produced similar rates of PTSD remission. Our finding of a large treatment effect for PTSD symptoms challenges the notion that exposure is a necessary component of PTSD and nightmare treatment, since SIP produced significant reductions in PTSD symptoms and nightmare frequency but involves very limited exposure to nightmare content.

Second, SIP incorporates an intervention that specifically targets insomnia and is heavily focused on regulation of erratic sleep schedules. Given the importance of homeostatic and circadian mechanisms on sleep and their role in sleep quality, it is our belief that behavioral sleep targets should be addressed first to provide the foundation for addressing other aspects of sleep disturbance (e.g., nightmares). Addressing nightmares without first targeting behaviors that serve to maintain insomnia is likely to impair the effectiveness of the nightmare intervention since sleep-disruptive behaviors may persist. However, in the absence of research showing a sequence effect, this assertion remains an empirical question.

As consistent with expectations for an intervention targeting sleep disturbance, SIP had the greatest impact on self-reported insomnia severity. SIP’s potent effect on PTSD symptoms, however, was a less expected and arguably, one of the more interesting findings of our study. The second largest treatment effect for SIP was found in the domain of PTSD symptoms (Cohen’s $d = 1.76$), and half of those in the intervention group scored in the subclinical PTSD screening range at post-intervention. This finding is consistent with the suggestion of some researchers that sleep plays a significant role in the development and/or maintenance of PTSD.

Krakov and colleagues have observed that many individuals with post-trauma symptoms, would prefer to address their insomnia first, then nightmares, and then PTSD, as applicable, and their observation is consistent with our clinical experience of PTSD patients seeking treatment for sleep disturbance. Acceptability of treatment is a highly relevant topic for a mental health condition for which avoidance is a primary cognitive feature. We did not assess the relative acceptability of sleep treatment versus PTSD treatment. However, if our clinical experience on this topic is based in fact, that those who are struggling with PTSD are more likely to seek and complete treatment for sleep disturbance, then it may be prudent to promote sleep disturbance interventions as a first-line PTSD treatment. Our evidence, along with the reports of other researchers, suggests that about half of participants could remit from PTSD following treatment for sleep disturbance.

As noted above, in spite of differences in approach, these interventions all facilitate a reduction in both nightmare frequency and PTSD symptoms. One explanation for this consistent finding across approaches is that ERRT, IRT, and SIP all promote an observer stance towards dream content. Several participants in our study reported a significant shift in nightmare frequency with the realization that they could alter the course of the dream while it was occurring. These participants described a shift from being part of the dream to being an observer of the dream. These reports are consistent with a shift towards metacognitive insight, as described by Teasdale, wherein “A distinction is made between metacognitive knowledge (knowing that thoughts are not necessarily always accurate) and metacognitive insight (experiencing thoughts as events in the field of awareness, rather than as direct readouts on reality)(pg. 146).” Recent findings of a study examining brain activity during self-related awareness tasks found that, by simply placing one’s attention on feelings and emotions one could modulate amygdala activation, thereby initiating the emotion regulation process. These authors propose that making oneself aware of emotions may provide a therapeutic distance that facilitates emotion regulation. Similarly, rather than promoting avoidance, IRT and similar approaches promote a stance of “observer” versus “experiencer” of affectively charged dream content.

**Limitations**

The findings of our study should be viewed in light of the limitations inherent to a pilot study. Namely, our sample size was very small, so we cannot be certain that the large effect sizes seen in this trial would be seen in a larger trial of SIP. Also, this study did not include an active control condition, and no follow-up data was collected. Thus, we cannot determine...
the effect of nonspecific factors, such as treatment expectancies and therapeutic alliance, and we cannot assure that our post-intervention outcomes translate to sustained beneficial effects. Davis et al.\textsuperscript{19} found that nightmare frequency improved from the post-intervention assessment to the follow-up assessments. Since we did not conduct a clinical assessment of PTSD using the CAPs at the 12-week assessment, we cannot assure that those screening negative for PTSD diagnosis following the intervention (PCL-M < 50) actually remitted from PTSD. Finally, we excluded veterans who screened positive for sleep apnea, since apnea produces its own sleep deprivation that would be unaddressed by our intervention. The inclusion of those with apnea would be further confounded by the observation of Krakow and colleagues\textsuperscript{42} that treatment of OSA alone resulted in subjective improvements in sleep, PTSD symptoms, and nightmares. However, since we did not assess participants with PSG, we cannot rule out the possibility that veterans with occult apnea were enrolled in our study. To assure the generalizability of SIP to the larger population of veterans with PTSD, a larger trial of SIP should include veterans with sleep apnea.

**Future Research**

All of those dropping out of the study were OEF/OIF veterans, and these veterans are also less likely to utilize VA clinical services.\textsuperscript{43} In future research, SIP will be tailored to assure that it is acceptable to and addresses the needs of our recently deployed veterans. With many younger veterans returning from deployment with trauma-related sleep disturbance, it will be critical to identify sleep interventions that are not only effective, but accessible. Sleep and trauma researchers should also attempt to determine the optimal sequence of treatment for those with PTSD. Should sleep disturbance be addressed before, after, or concurrent with PTSD treatment, and what are the characteristics of those who remit from PTSD following a trauma-focused sleep intervention versus those who do not remit?

In spite of the limitations of our study, the findings demonstrate that an intervention targeting both insomnia and nightmares produces large short-term effects. The collective findings of our research along with those of other sleep and trauma researchers suggest that the treatment of sleep disturbance produces significant reductions in PTSD symptoms, in addition to the amelioration of sleep disturbance. Thus, the field of behavioral sleep medicine is poised to extend the reach of our trauma-specific interventions beyond sleep clinics and into mental health clinics through the provision of clinical services and mental health provider training.

**REFERENCES**

ACKNOWLEDGMENTS

The first author was funded by a Department of Veterans Affairs HSR&D Career Development Award CDA 09-218. The authors would like to thank Maren Olsen, Ph.D., and Daniel Almirall, Ph.D., for their statistical support in preparing this manuscript. This material is based upon work performed at the Durham VA Medical Center and supported by the Institute for Medical Research and the Department of Veterans Affairs, Veterans Health Administration, Office of Research and Development.

SUBMISSION & CORRESPONDENCE INFORMATION

Submitted for publication June, 2010
Submitted in final revised form September, 2010
Accepted for publication September, 2010
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DISCLOSURE STATEMENT

The views expressed in this article are those of the authors and do not necessarily reflect the position or policy of the Department of Veterans Affairs or the United States Government. This was not an industry supported study. Dr. Edinger has received research support from Philips Respironics and has consulted for Philips Respironics and Kingsdown, Inc. The other authors have indicated no financial conflicts of interest.