Feasibility study of unattended polysomnography in medical intensive care unit patients

Melissa P. Knauert, MD, PhD\textsuperscript{a,}\textsuperscript{*}, H. Klar Yaggi, MD, MPH\textsuperscript{a}, Nancy S. Redeker, PhD, RN\textsuperscript{b}, Terrence E. Murphy, PhD\textsuperscript{c}, Katy L. Araujo, MPH\textsuperscript{c}, Margaret A. Pisani, MD, MPH\textsuperscript{a}

\textsuperscript{a}Department of Internal Medicine, Section of Pulmonary, Critical Care, and Sleep Medicine, Yale University School of Medicine, 333 Cedar Street, P.O. Box 208057, New Haven, CT 06520-8057, USA
\textsuperscript{b}Yale School of Nursing, Rm 20508, Yale University West Campus, P.O. Box 27399, West Haven, CT 06536, USA
\textsuperscript{c}Department of Internal Medicine, Section of Geriatrics, Yale University School of Medicine, 333 Cedar Street, P.O. Box 208056, New Haven, CT 06520, USA

A R T I C L E   I N F O

Article history:
Received 19 March 2014
Received in revised form 7 June 2014
Accepted 9 June 2014
Available online 12 July 2014

Keywords:
Sleep deprivation
Sleep fragmentation
Polysomnography
Critical care
Feasibility study

A B S T R A C T

Objectives: To evaluate the feasibility of using unattended, portable polysomnography (PSG) to measure sleep among patients in the medical intensive care unit (MICU).

Background: Accurate measurement of sleep is critical to studies of MICU sleep deprivation. Although PSG is the gold standard, there is limited data regarding the feasibility of utilizing unattended, portable PSG modalities in the MICU.

Methods: MICU based observational pilot study. We conducted unattended, 24-h PSG studies in 29 patients. Indicators of feasibility included attainment of electroencephalography data sufficient to determine sleep stage, sleep efficiency, and arousal indices.

Results: Electroencephalography data were not affected by electrical interference and were of interpretable quality in 27/29 (93%) of patients. Overnight sleep efficiency was 48% reflecting a mean overnight sleep duration of 3.7 h.

Conclusions: Unattended, portable PSG produces high quality sleep data in the MICU and can facilitate investigation of sleep deprivation among critically ill patients. Patient sleep was short and highly fragmented.

© 2014 Elsevier Inc. All rights reserved.

Introduction

Sleep deprivation, characterized by short sleep duration and poor sleep continuity (fragmentation), is virtually universal in critically ill patients.\textsuperscript{1} A variety of environmental, patient care and physiologic factors influence sleep deprivation. Noise disrupts patient sleep,\textsuperscript{2} and patients experience high levels of overnight in-room activity.\textsuperscript{3,5} Furthermore, patients undergoing mechanical ventilation and accompanying continuous sedation suffer particular decrements in sleep quality.\textsuperscript{4,5} Because sleep deprived patients often demonstrate behaviors that mimic intensive care unit (ICU) delirium,\textsuperscript{3} there is concern that sleep deprivation potentiates ICU delirium,\textsuperscript{6} and therefore contributes to delirium-related morbidity and mortality.\textsuperscript{7,11} Additionally, complete sleep deprivation and selective rapid eye movement (REM) sleep and slow wave sleep deprivation are associated with immune system dysfunction.\textsuperscript{14,15} Animal models and studies in other patient populations suggest links to other negative outcomes including derangements in glucose metabolism,\textsuperscript{16,17} adverse cardiovascular events,\textsuperscript{18} cognitive dysfunction,\textsuperscript{19,20} and all-cause mortality.\textsuperscript{21}

Polysomnography (PSG) is the gold standard of sleep measurement and allows full delineation of sleep duration, continuity, and architecture, characterized by sleep stages and electrophysiological arousals. Reports of PSG use in the ICU have focused primarily on mechanically ventilated patients.\textsuperscript{8,22,23} Studies of non-ventilated patients have most commonly included healthy volunteers in

\textbf{List of abbreviations:} AASM, American Academy of Sleep Medicine; APACHE II, Acute Physiology and Chronic Health Evaluation II; CAM-ICU, Confusion assessment method for the ICU; ECG, electrocardiogram; EEG, electroencephalography; EMG, electromyogram; EOG, electrooculogram; Hz, hertz; ICU, intensive care unit (specialty not designated); MICU, medical intensive care unit; NREM, non-rapid eye sleep; PSG, polysomnography; RASS, Richmond agitation sedation scale; REM, rapid eye movement sleep; RPSGT, Board certified sleep technician; SD, standard deviation.

Sources of funding: This work was supported by a grant to Dr. Pisani (5R21NR11066), by a grant to Drs. Knauert, Yaggi and Redeker (5P20NR014126), and by a grant to Dr. Murphy and Ms. Araujo (P30AG21342).

The authors declare that they have no conflict of interests.

\textsuperscript{*} Corresponding author. Tel.: +1 203 785 4163.
\textit{E-mail address:} melissa.knauert@yale.edu (M.P. Knauert).

\textsuperscript{1} Sources of funding: This work was supported by a grant to Dr. Pisani (5R21NR11066), by a grant to Drs. Knauert, Yaggi and Redeker (5P20NR014126), and by a grant to Dr. Murphy and Ms. Araujo (P30AG21342).
simulated ICU environments. Barriers such as large amounts of technician time, concerns about lead displacement, electrical interference, and difficulties with scoring sleep in the ICU have been noted by previous investigators but not rigorously studied.

Existing PSG studies demonstrate that the sleep of critically ill patients is severely disrupted. ICU patients have more difficulty falling asleep, more difficulty progressing through sleep stages, more arousals, decreased REM sleep, and decreased N3 sleep. In addition, sleep is distributed throughout the 24-h day rather than consolidated during the night, as during normal circadian cycling. Several groups have identified a subgroup of ICU patients with atypical sleep patterns of unknown clinical significance. Atypical sleep's most prominent characteristic is electroencephalography (EEG) tracings consistent with sleep but lacking sleep spindles, K complexes and other markers of specific non-REM (NREM) sleep stages. REM sleep has also been identified as diminished in this population.

Given the need for PSG measures of ICU sleep and the limited data regarding barriers to data acquisition and scoring, the objective of this study was to establish the feasibility of monitoring the sleep of medical ICU (MICU) patients via unattended portable PSG (type 2 device) for a 24 h period. Fragmentation of ICU patient sleep across the 24 h day led us to include both day and night recording periods. Technical feasibility outcomes included the ability to obtain EEG data of sufficient quality to determine sleep stages, sleep efficiency, and arousals from sleep. Acceptability outcomes included patient, family and staff acceptance of PSG, and a review of the reasons given for premature discontinuation.

Methods

Study design and setting

This was an observational, cross-sectional study. We obtained approval for the study from the Institutional Review Board and informed consent was obtained from all participants. The study was conducted at a 1500 bed academic teaching hospital in New England with a 38 bed closed MICU staffed by university faculty physicians. The MICU is new construction (completed in 2010) with all private rooms separated by solid walls and glass sliding doors. The unit is rectangular in design with an administrative desk at the single main entrance. The central core of the rectangle is comprised of enclosed supply and conference rooms. There is no central nursing station. Most clinical staff work 7:00 AM to 7:00 PM shifts. Lab work, wound care, bathing and routine radiology most typically occur overnight (specifically within the 10:00 PM to 6:00 AM period).

Inclusion and exclusion criteria, patient characteristics

We included English-speaking patients older than 18 years of age who were admitted to the MICU for less than 72 h and who were expected to stay for at least an additional 24 h at the time of screening. Exclusion criteria included terminal illness, receipt of comfort care only, coma or deep sedation (Richmond Agitation Sedation Scale (RASS) score of −4 to −5), patient inability to consent and without an identifiable surrogate, anticipated procedure requiring sterile access to the head or neck during the PSG recording time, anticipated procedure requiring movement of patient out of the MICU during the PSG recording time, severe agitation (agitated to touch, hallucinations, uncontrolled pain, violent behavior), and anatomic contraindication to PSG (including infections of the head or neck, recent surgery of the head or neck, recent trauma to the head or neck). We did not exclude patients who were mechanically ventilated and did not use a severity of illness scoring system cut-point for study exclusion.

Medication usage was abstracted for the 48 h prior to PSG initiation and during the PSG study. All medications, regardless of administration route, from the following classes were recorded: sleep aids, benzodiazepines, opioids, and vasoactive medications. Home medication use was included for study subjects who had their PSG initiated within 48 h of hospital admission. If an entire day at home fell within the 48 h prior to PSG initiation, all home medications were presumed to be taken unless this was explicitly contradicted in the chart. If a portion of a day fell within the 48 h prior to PSG initiation, hospital arrival time was used to estimate what medications were likely taken prior to admission. Daily medications were presumed to be taken in the morning unless prescribed with a per bedtime designation. Twice daily medications were presumed to be taken in the morning and evening and thrice daily medications were presumed to be taken in the morning, midday and evening.

Study procedures

Patients or their surrogates gave verbal consent for study participation. A waiver of signed consent was granted by the Institutional Review Board. Surrogates were approached for consent in the event of patient delirium (positive Confusion Assessment Method for the ICU (CAM-ICU)) status,40,41 patient sedation (RASS −1 to −5) and/or inability of the patient to communicate with study staff. No compensation was offered for study participation. Reasons for ineligibility were recorded. Patient demographic and medical data were collected for enrolled and non-enrolled patients to assess for enrollment bias secondary to patient age, sex, diagnosis, severity of illness or ventilator status. The Acute Physiology and Chronic Health Evaluation II (APACHE II)42 scoring algorithm was utilized for estimation of illness severity. Delirium status was also monitored on the day of enrollment and during PSG recording via CAM-ICU40,41 and chart review.43

Polysomnography

Patients underwent unattended, 24-h PSG obtained with the SAFIRO Portable Data Acquisition System (Compumedics, Abbotsford, VIC, Australia) in the setting of otherwise usual MICU care. PSG studies were initiated in the evening and terminated at or before 24 h. Leads were applied by a board certified sleep technician (RPSGT) familiar with the MICU clinical setting. Leads included 6 EEG channels (C3, C4, F3, F4, O1, O2) referenced to the contralateral mastoid. A chin electromyogram (EMG), right and left electrooculograms (EOC), and electrocardiogram (ECC) were also recorded. EEG signals were amplified, recorded at 200 Hz sampling frequency and filtered (0.5–70 Hz). The signals were recorded on the SAFIRO device and later transferred to a laptop computer with Profusion 2 software (Compumedics, Abbotsford, VIC, Australia). The studies were not attended by the technician once the leads were applied and recordings were started. The MICU nurses responsible for caring for the patients were instructed on lead removal and safe equipment shut down in the event of transfer or clinical need. They were not instructed to replace leads in the event of disconnection.

Information regarding duration of the PSG study and reasons for early termination of the PSG study were collected. PSG data were scored by RPSGTs according to standard American Academy of Sleep Medicine (AASM) protocols. The technologist scoring this data participated in the AASM inter-rater reliability program and maintained a 93% agreement with AASM epochs over the most recent eight month period reviewed at the time of the study. Individual epochs were scored as wake, sleep stage N1, sleep stage N2, sleep stage N3 or REM sleep. If a determination could not be made for a particular epoch, the epoch was marked “unsure,” and if the epoch could be identified as sleep, but the stage of sleep was unclear, the epoch was marked “sleep” (no stage). EEG scoring and PSG tracings...
were reviewed by sleep medicine physicians (HKY and MPK). Sleep data obtained during PSG were analyzed for sleep duration, diurnal sleep timing, sleep stage distribution, arousal indices and the presence or absence of atypical sleep features. Patients were considered to have atypical sleep following the definitions of Cooper et al. Briefly, these patients had EEG consistent with stage N2 sleep but lacking sleep spindles and K complexes. Definitions for sleep related terminology are provided in Table 1.

**Data analysis**

Data analyses were performed with SAS software V9.3 (SAS Institute, North Carolina). Continuous variables were expressed as a mean with standard deviation (SD) or median with interquartile range. Frequencies and percentages were used to express categorical data. Differences in frequencies or percentages between groups were tested with a chi-square statistic; differences in means were tested with an unpaired Student’s t-test; and differences in medians were tested with the Wilcoxon Two Sample statistic. APACHE II score was calculated according to published algorithms.

PSG studies with a length of less than 4 h were examined for feasibility metrics but were excluded from analysis of sleep architecture. For calculation of sleep efficiency, stage proportions, and arousal indices, the whole study was considered to be from the first to last study epoch. Overnight was considered to be from 10:00 PM to 6:00 AM and all other hours were considered as daytime. For each individual patient, sleep efficiency was calculated by dividing total sleep time by total time available for sleep for a given period (whole study, overnight, or daytime). Sleep stage proportions were similarly calculated by dividing the amount of a given stage of sleep by the total sleep time in a given period. Arousal indices were calculated by dividing the total number of arousals in either REM or NREM sleep by the total amount of REM or NREM sleep in hours. A mean was then calculated across the study population for each sleep variable included in the sleep architecture analysis.

**Results**

**Patient characteristics**

We screened 275 patients, of whom 181 patients were ineligible, yielding 94 eligible patients (see Fig. 1). Ineligible status was due to a variety of factors; lack of identifiable surrogate (56), patient was comatose (32) and patient expected to die within the next 24 h (26) were dominant reasons. Of the 62 eligible, non-enrolled patients, 20 patients were not approached because consent was not granted by a member of the clinical team; the other 42 patients declined to participate. Thirty-two eligible patients were enrolled. Two patients who were initially enrolled subsequently declined further participation once exposed to the lead application process. One patient deteriorated clinically, and the family requested withdrawal from the study. Ultimately 29 patients had PSG studies initiated; feasibility analysis was based on this cohort.

Baseline demographic, medical and severity of illness characteristics of the enrolled patients for whom PSG (N = 29) was attempted are described in Table 2. Patients had a mean age of 59.2 years and average length of MICU stay of 10.9 days. Reasons for ICU admission were diverse and dominated by sepsis and respiratory failure. Mean APACHE II score was 13.9 with 24% of patients dependent on mechanical ventilation. Comparison of enrolled and non-enrolled MICU patients reveals no significant differences in age, gender, race, reason for ICU admission, severity of illness or mechanical ventilation status. Furthermore, the patients with PSG studies greater than 4 h in length (n = 23) and whose PSGs were ultimately included in the sleep architecture analyses did not differ significantly in age, gender, race, reason for ICU admission, severity of illness or mechanical ventilation status compared to the general enrollment group (N = 29) or the non-enrolled patients (data not shown).

Patient medication use included drugs that influence sleep (see Table 3). Thirty-five percent (10/29) of all enrolled patients received benzodiazepines; sixty-six percent (19/29) received opioids; seven percent (2/29) received sleep aids. Thirty-one percent (9/29) of patients received none of these three classes of medication. No study patients received propofol or dexmedetomidine.

**PSG data quality**

Unattended PSG sleep data were interpretable using standard AASM criteria in the 23 studies included in the sleep architecture analysis; these studies were selected on the basis of length (greater than 4 h). The sleep data obtained from 4 additional studies were interpretable, but were excluded from sleep architecture analysis on the basis of short duration (less than 4 h). Two studies had no interpretable data. There was no evidence of electrical (60 Hz)
interference from the ICU equipment; devices known to be on or near patients during PSG recording included telemetry nodes and leads, oximetry nodes and leads, mechanical ventilators, mobile computer units (utilized for bedside recording into the electronic medical record), sequential compression devices, and hemodialysis units (non-continuous).

The proportion of epochs in all interpreted studies that was scored as unsure was 9.5%. Re-evaluation of these studies, using the criteria of Cooper et al.\(^\text{35}\) and Drouot et al.\(^\text{30}\) for atypical ICU sleep revealed that 9 of 26 (35%) patients with interpretable EEG and at least one epoch of sleep present during their study had atypical critical care sleep patterns; of the 27 patients with interpretable sleep data, 1 patient with a short PSG had only wake recorded and could therefore not be assessed for atypical sleep. Fig. 2 presents a sample EEG tracing of atypical sleep. Based on CAM-ICU and chart review, 4 of 9 (44%) patients with atypical sleep had clinical evidence of delirium as compared to 3 of 17 (18%) of patients with typical sleep; this difference did not reach statistical significance.

Atypical sleepers received benzodiazepines 11% (1/9) of the time, opioids 56% (5/9) of the time and on sleep aids 11% (1/9) of the time. Forty-four percent (4/9) of atypical sleepers received none of these three classes of medication. Typical sleep patients received benzodiazepines 50% (7/14) of time, opioids 71% (10/14) and sleep aids 7% of the time (1/14). Twenty-one percent (3/14) of typical sleepers received none of these three classes of medication (see Table 3).

For typical sleep patients, repeat analysis of epochs marked as "unsure" by the sleep technician allowed classification of most epochs as either wake or stage N1 sleep. The majority of the uncertainty reflected difficulty in identifying the transition from wake to sleep in patients with frequent variation in sleep wake status (i.e. micro-sleeps). Following review of all sleep studies by sleep medicine physicians (MPK and HKY), and, considering atypical sleep, the proportional amount of scoring uncertainty was reduced to 0.3%.

**PSG study duration**

Nocturnal recordings of at least 4 h in length were accomplished for 23 of 29 patients. For all initiated studies, median sleep study duration was 15.1 h (Table 4A). Reasons for unattended PSG discontinuation are delineated in Table 4B. Patient request for discontinuation was the most common reason for incomplete studies with 9 of 29 patients (31%) requesting leads to be removed before completing 24 h of monitoring; typically these patients requested lead removal in the early morning upon “waking from sleep.” Qualitative comments from patients requesting lead

---

**Fig. 1.** Patient enrollment, screening, and eligibility flow diagram. "✖️" with number indicates patients excluded from PSG sleep architecture analysis. h indicates length of study in hours; PT indicates patient; PSG indicates polysomnogram.
Table 2
Clinical and demographic characteristics of enrolled study participants.

<table>
<thead>
<tr>
<th>Enrolled</th>
<th>N = 29</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline demographics</td>
<td></td>
</tr>
<tr>
<td>Age in years: mean (SD)</td>
<td>59.2 (17.8)</td>
</tr>
<tr>
<td>Male gender: n (%)</td>
<td>19 (66)</td>
</tr>
<tr>
<td>Race, non-white: n (%)</td>
<td>12 (41)</td>
</tr>
<tr>
<td>Mechanical ventilation: n (%)</td>
<td>33 (11)</td>
</tr>
<tr>
<td>ICU length of stay in days: mean (SD)</td>
<td>10.9 (7.7)</td>
</tr>
<tr>
<td>Time in sleep study in days: mean (SD)</td>
<td>3.6 (2.8)</td>
</tr>
</tbody>
</table>

Reason for ICU admission: n (%)

- Infection, sepsis: 11 (38)
- Respiratory failure: 5 (17)
- Heart failure: 1 (3.5)
- Pulmonary embolism: 1 (3.5)
- Gastrointestinal bleed: 1 (3.5)
- Liver disease: 1 (3.5)
- Acute kidney injury: 2 (7)
- Neurologic injury: 2 (7)
- Other: 5 (17)

Severity of illness parameters

- APACHE II score: 20.7 (7)
- Mechanical ventilation: 7 (24)
- Delirium, n (%) | 10 (34)

equipment removal reflected patient discomfort with leads and a desire to move from bed to chair. Two patients requested lead removal at or before 4 h of data acquisition which resulted in exclusion of their PSG data from sleep architecture analysis; both of these patients named lead discomfort and inability to sleep as the reason for their request. Patient transfer out of the MICU to the floor was also common (8 of 29 studies, 28%).

Patients who were mechanically ventilated (n = 7) had a median (interquartile range) study duration in hours of 16.9 (11.7). There was no significant difference in study duration compared with non-ventilated patients. As might be expected, these patients were more severely ill than the non-ventilated study sample (mean APACHE II score = 20.7, P = 0.01). Mechanically ventilated patients requested discontinuation (1 of 7, 14%) and were transferred (1 of 7, 14%) less frequently than the general study population (see above). There were trends suggesting that the patients in the most severely ill quartile (APACHE scores 19–33) had longer studies with a mean (SD) duration in hours of 14.0 (6.3) versus those in the least severely ill quartile (APACHE scores 2–8), with mean (SD) duration in hours of 7.9 (5.7) (P = 0.09).

Technical limitations

Technical difficulties occurred in 8 of 29 PSG studies. Technical difficulties were significant enough to be the primary reason for PSG discontinuation for 5 of them. In 3 cases, these technical problems led to complete PSG data exclusion. One excluded patient had significant scalp edema and although leads remained in place for the full 24-h, EEG recording quality was severely limited and therefore uninterpretable. A second patient experienced equipment failure with no data recorded on the Safiro device; leads remained in place without difficulty throughout this study. Unexplained data truncation occurred in 3 of 29 patients, although in only one case did this cause significant shortening of the study (PSG length 12 h or less) and resultant data exclusion. Unintended lead removal occurred in 3 of 29 patients (10%) and, in two cases, this caused significant shortening of the study (PSG length 12 h or less). For the three patients with lead removal, continuous nebulization treatment and patient delirium were factors that contributed to difficulty with lead maintenance.

Patient sleep characteristics

For all patients, the overall mean (SD) sleep efficiency was 38.0% (16.6) for the entire period of PSG recording. Analysis of sleep between 10:00 PM and 6:00 AM revealed an overnight mean (SD) sleep efficiency of 48.8% (20.5). During daytime recording hours (6:00 AM to 10:00 PM), patients were asleep 29.2% (SD = 23.3) of the time. Mean (SD) wake after sleep onset in the overnight period was 155 (85.6) minutes. Representative 10:00 PM to 6:00 AM hypnograms reveal the highly fragmented overnight sleep of study patients (Fig. 3). Review of the sleep architecture demonstrated a paucity of N3 slow wave sleep (3.9% of total sleep time) and REM sleep (10.5% of total sleep time). Stage N1 sleep comprised 25.8% of all sleep and stage N2 sleep comprised 59.7% of all sleep. Stage distribution did not vary significantly in overnight versus daytime sleep; there was a trend toward increased REM proportion during daytime versus overnight hours. Patients experienced frequent arousals during their sleep with an overall NREM arousal index of 30.4 per hour and a REM arousal index of 16.4 per hour. This rate of arousal did not vary significantly between overnight and daytime sleep. Table 5A and B presents sleep parameters for typical and atypical patient subsets. Patients with typical and atypical sleep patterns had similar sleep efficiencies, sleep architecture and arousal indices.

Discussion

We have demonstrated that it is technically feasible to obtain high quality unattended polysomnographic data among critically ill patients. However, several patients tolerated the PSG procedure poorly. ICU patient tolerance of PSG has not been reported in the literature and may be related to the inclusion of non-ventilated patients in our study population. We felt this was an important group to study as this group may be more easily targeted for sleep improvement interventions.

Readily interpretable sleep EEG data were recorded in 93% (27/ 29) patients. Consistent with past studies, patient sleep in the MICU was highly fragmented and was distributed across day and night time periods.38,47 Sleep efficiencies overnight were low, and there was an abnormally high amount of sleep during the daytime period. Total sleep time of 5.6 h indicates generalized sleep deprivation, and an evaluation of the distribution of the sleep stages indicates N3 and REM specific sleep deprivation. Additionally, approximately one third of patients had evidence of atypical sleep.

Technical issues were encountered during our unattended PSG studies. Data truncation and data loss were the most significant technical issues and did significantly shorten studies (PSG time less than 12 h). This likely could have been avoided had the studies been attended. Lead instability contributed to shortening of several studies, though this caused significant shortening (study length less than 12 h) of the study in only 2 cases. Despite data and lead issues, ICU-based use of unattended PSG conveys a cost and personnel savings in comparison to fully attended monitoring, and, therefore, is a more efficient strategy overall. In future studies, nurse training could be performed to minimize some but not all of the lead...
complications. There was no ICU equipment interference with our EEG recordings, despite previous reports of this problem in the literature. Overall, we judge unattended PSG in the MICU to be technically feasible.

Notably, patient, family and caregiver tolerance or perceived tolerance of PSG is poor. This is reflected in the large number of patients deemed “inappropriate” by a member of their caregiving team, the low screen to enrollment ratio and the frequent request by patients to have their studies discontinued. Patients specifically described the leads as limiting mobility and their ability to sleep. Severity of illness did not appear to influence enrollment, and the more severely ill patients tended to have longer unattended PSG recording times. The fact that these patients were unlikely to be transferred and/or possibly unable to request study discontinuation likely contributed to the longer recording times. Patients undergoing mechanical ventilation were less likely to request lead discontinuation than non-ventilated patients perhaps due to the severity of their illness and/or related to their inability to communicate.

Half of our patients had an APACHE II score less than or equal to 14 (estimated hospital mortality 15% or less). Among these patients with a lower severity of illness, turnover of patients in the MICU played a role in PSG study length. About one third of patients with a lower severity of illness, turnover of patients in the MICU played a role in PSG study length. About one third of patients with a lower severity of illness, turnover of patients in the MICU played a role in PSG study length. About one third of patients with a lower severity of illness, turnover of patients in the MICU played a role in PSG study length. About one third of patients with a lower severity of illness, turnover of patients in the MICU played a role in PSG study length.

Table 4

<table>
<thead>
<tr>
<th>Reason for Discontinuation</th>
<th>All Patients</th>
<th>Ventilated Patients</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. PSG recording time in hours</td>
<td>n = 28&lt;sup&gt;a&lt;/sup&gt;</td>
<td>n = 7</td>
</tr>
<tr>
<td>Minimum value</td>
<td>1.4</td>
<td>6.3</td>
</tr>
<tr>
<td>1st quartile</td>
<td>11.0</td>
<td>9.9</td>
</tr>
<tr>
<td>Median</td>
<td>15.1</td>
<td>16.9</td>
</tr>
<tr>
<td>3rd quartile</td>
<td>19.0</td>
<td>21.6</td>
</tr>
<tr>
<td>Maximum value</td>
<td>24.0</td>
<td>21.8</td>
</tr>
<tr>
<td>B. PSG reason for discontinuation prior to 24 h goal</td>
<td>n = 28&lt;sup&gt;a&lt;/sup&gt;</td>
<td>n = 7</td>
</tr>
<tr>
<td>Patient request</td>
<td>9 (31%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Patient transfer</td>
<td>8 (28%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>PSG technical failure</td>
<td>5 (17%)</td>
<td>3 (43%)</td>
</tr>
<tr>
<td>Other</td>
<td>3 (10%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>Patient condition deterioration</td>
<td>2 (7%)</td>
<td>1 (14%)</td>
</tr>
<tr>
<td>24 h completed</td>
<td>2 (7%)</td>
<td>0 (0%)</td>
</tr>
</tbody>
</table>

<sup>a</sup> No data recorded on one patient; length of study unknown.

Fig. 2. Electroencephalography tracing from an atypical sleep patient. LOC and ROC indicate left and right electrooculogram respectively. C3, C4, F3, F4, O1, O2 indicate EEG lead locations. EMG indicates electromyogram and ECG indicates electrocardiogram.
ventilated counterparts. They are similarly at risk for ICU delirium and its downstream morbidity and mortality. In addition it may be easier to improve upon their sleep quality without the significant barrier of overcoming mechanical ventilation related sleep disturbance. A challenge to further study of these patients will be improving patient, family and caregiver acceptance of PSG and patient tolerance of the PSG lead application and monitoring.

Study limitations

This study was focused upon assessment of the technical feasibility of unattended PSG. Acceptability by patients, family and clinical staff emerged as a limitation to PSG use. The somewhat low screen to enrollment rate reflected a significant lack of acceptance by clinical staff who indicated that their patients were not appropriate for study participation. In addition, patient discomfort and patient transfer during PSG recording time significantly truncated many studies. As these were not anticipated barriers, they were not investigated with formal instruments. It is somewhat reassuring that patient demographics, reason for ICU admission, severity of illness (APACHE II) and ventilator status were similar between enrolled and non-enrolled patient groups. It is notable that less severely ill patients and non-ventilated patients had generally shorter recording times which reflected a higher rate of patient transfer and the ability to communicate and therefore request lead removal. Despite these limitations, our study demonstrates sleep architecture and arousal indices in-line with other ICU studies. Our primary goal of investigating the technical feasibility of unattended PSG use was accomplished.

Implications for practice

The findings of this study, in conjunction with other ICU studies of sleep deprivation, suggest that patient sleep in the ICU is profoundly disrupted. Clinical providers need to invest greater attention to sleep promotion during day and night periods. Night staff should avoid unnecessary overnight disturbance of patients, and day staff should encourage wakefulness during the daytime period. Less severely ill and non-ventilated ICU patients experience severe sleep loss, along with their more ill and/or ventilated counterparts, and may be an ideal target population for sleep improvement interventions.
Implications for future research

PSG provides unique and important data to ICU sleep deprivation studies. Future explorations of atypical sleep would require sleep monitoring that includes EEG leads. The successful use of an unattended PSG, inclusive of EEG without electrical interference or severe data loss, indicates that this is an accurate, cost-effective modality for use in ICU research studies. There are issues with patient, family and staff acceptability that pose a barrier to patient enrollment.

Conclusions

Unattended PSG with full sleep EEG montage is a technically feasible methodology for the study of sleep in the ICU. Technical limitations were acceptably low and data produced from these studies were generally of high quality. Patient intolerance of the PSG procedure was largely due to discomfort with leads. MICU sleep was highly disrupted with prominent loss of REM and N3 sleep. Critically ill patients had deranged circadian sleep patterns with abnormally high amounts of sleep occurring during the day. About one third of patients demonstrated atypical sleep. Recognition of this unusual sleep pattern clarified prior difficulties in scoring the sleep of ICU patients.

Acknowledgments

We appreciate the assistance of our RPSGT colleagues Elizabeth Lowe, Vincent McClain and Rebecca Khozein who executed the polysomnography studies.

References