Circadian rhythm in the assessment of postconcussion insomnia: a cross-sectional observational study

Dora M. Zalai MD PhD, Todd A. Girard PhD, Michael D. Cusimano MD PhD, Colin M. Shapiro MBBCh PhD

Abstract

Background: Insomnia is a major predictor of adverse outcomes in mild traumatic brain injury (mTBI), including concussion; although insomnia symptoms may be due to various sleep disorders, those related to circadian rhythm sleep–wake disorders (CRSWDs) require specific assessment and treatment. The objective of the current study was to determine the prevalence of CRSWD in a sample of treatment-seeking people with chronic insomnia symptoms after an mTBI.

Methods: Participants aged 17–65 years who had experienced an mTBI and reported chronic insomnia were recruited from diverse community clinics in Ontario 3–24 months after their injury to participate in this cross-sectional observational study. Potential participants were screened by both telephone and intake interview. Exclusion criteria were alcohol or substance use disorders, preexisting brain disorder or previous neurosurgery, recent travel across more than 2 time zones or shift work. Assessments included a clinical interview, questionnaires, 2 weeks of actigraphy and a sleep diary, and a dim-light melatonin onset test. The main outcome measure was the proportion of patients with CRSWDs.

Results: Of the 50 participants (32 [64%] female; median age 39.5 yr), 13 (26% [standard deviation 12%]) had an CRSWD. The most common circadian diagnosis was delayed sleep–wake phase disorder (10 participants [20%]).

Interpretation: The prevalence of CRSWDs may be exceptionally high among people with chronic insomnia symptoms following mTBI. Proper detection and treatment of CRSWDs in this population is essential to facilitate recovery. The findings emphasize the relevance of a diagnostic circadian assessment in patients with mTBI presenting with chronic insomnia symptoms.

Traumatic brain injury (TBI) is the leading cause of disability among young adults, and the incidence is increasing in Canada.1,2 Mild TBI (mTBI), including concussion, accounts for the majority (70%–90%) of treated TBI cases and has the largest contribution to injury-related disability.3 Early detection and effective management of medical conditions that hinder recovery could prevent disability and substantially reduce societal costs.

In this regard, sleep disruptions are key modifiable targets that magnify the sequelae of mTBI.1 Insomnia — difficulties falling or staying asleep, or early-morning final awakening — is the most common persistent sleep symptom after mTBI1–6 and is particularly prevalent following mTBI compared to more severe forms.4 Moreover, insomnia after concussion worsens fatigue, pain, cognition and mood, and predicts poor overall prognosis for recovery, greater disability and mental disorders.7–11

Given that disorders of sleep and wakefulness can be treated effectively, treatment holds the promise of improving the management of persistent postconcussive symptoms and hastening recovery from mTBI. Recent reports4,12,13 emphasize the need for multimodal sleep and circadian assessments; however, these are not common practice, and research remains limited.

Standard assessment of insomnia relies on patients’ subjective report, and the recommended first-line treatment is cognitive behavioural therapy.14,15 Importantly, however, circadian rhythm sleep–wake disorders (CRSWDs) that may underlie symptoms of insomnia require fundamentally different assessment and treatment (Appendix 1, available at www.cmajopen.ca/content/8/1/E142/suppl/DC1). These disorders arise if there is a disruption of the endogenous circadian system or a misalignment between the internal circadian rhythm and the external environment. The best biologic marker of CRSWDs is abnormal dim-light onset of melatonin production, and the behavioural diagnostic feature is that the sleep phase is markedly delayed, advanced or irregular relative to environmental time and social norms16 (Appendix 1).

Rates and mechanisms of CRSWDs in mTBI are not well characterized, but evidence indicates that brain trauma can disrupt systems that regulate circadian rhythms, such as suprachiasmatic nucleus function, gene expression, neural plasticity,
and melatonin production and transmission. For instance, mTBI may cause pineal gland injury and disrupt circadian regulation via its effect on melatonin secretion.

To our knowledge, only 1 clinical study has examined diagnostic circadian assessment in patients with mTBI. A high prevalence of CRSWDs was found among participants.

Given that CRSWDs require specific circadian assessment and treatment, it is pertinent to establish whether patients with mTBI and insomnia symptoms constitute one of the few clinical groups in which CRSWDs are prevalent. If so, circadian assessment and treatment becomes an important part of chronic insomnia investigation and management following mTBI.

The objective of the current study was to determine the prevalence of CRSWDs (according to standard diagnostic criteria using evidence-based comprehensive assessment including actigraphy and dim-light melatonin onset) in a sample of treatment-seeking people with chronic insomnia following mTBI. Our hypothesis was that CRSWDs are more common in this clinical group than among people who seek treatment for chronic insomnia in sleep clinics.

Methods

This was a cross-sectional observational study. Data were collected at a sleep (Sleep and Alertness Clinic) and a melatonin testing laboratory in Toronto in 2016 and 2017, and were analyzed in 2018.

Participants

Information about the study was distributed widely to hospitals and community clinics in southern Ontario (primarily the Greater Toronto Area) and lower regions of northern Ontario, posted on brain injury and sleep clinic websites, and disseminated by word of mouth. Referral sources included a range of clinics and specialists.

Participants were included if they had had an mTBI (Glasgow Coma Scale score ≥ 13 at the time of injury) 3–24 months before the screening assessment, had chronic insomnia symptoms that started or amplified markedly after the injury, and were aged 17–65 years. If a participant had multiple concussions, we calculated time elapsed since the injury, and were aged 17–65 years. If a participant had had a concussion that preceded the onset of the sleep problem (3–24 mo). Participants were excluded if they had alcohol or other substance use problems within 3 months before enrolment based on the criteria of the fifth edition of the Diagnostic and Statistical Manual of Mental Disorders (DSM-5), needed immediate psychiatric help based on in-person assessment, had preexisting brain disorders or had had neurosurgery, had travelled across more than 2 time zones less than 3 weeks before the assessment or had a shift work schedule.

Procedures

Prospective participants were screened initially with a telephone interview (5–10 min). The telephone screening assessment included questions about the presence of insomnia symptoms, history of brain injury, preexisting brain disorder, age, substance use, travel across time zones and shift work history. Those who passed the telephone screen participated in an in-person assessment that included a standard interview and completion of questionnaires (90–120 min). When family members accompanied participants, they were present during the interview and provided collateral information with participants’ consent.

In the 12–14 days after their enrolment, participants kept a sleep diary and wore a Philips Respironics Actiwatch on their nondominant wrist. They participated in a dim-light melatonin onset test 12–14 days after their initial interview.

Assessment tools and methods

Insomnia assessment

The Duke Structured Interview for Sleep Disorders assesses sleep disorders according to clinical and research diagnostic criteria. We used an updated version of this tool in which the insomnia section has been modified to match the new DSM-5 criteria to select participants with chronic insomnia.

The Insomnia Severity Index measures the subjective severity of insomnia symptoms. The degree of insomnia severity is determined by the summary scores. A summary score of 11 has been recommended as a cut-off for screening for clinical insomnia research and was used as cut-off for inclusion in the current study.

Assessment of sleep pattern

Wrist actigraphy is a recommended objective diagnostic assessment tool for CRSWDs. Participants wore a Philips Respironics Actiwatch 2 on their nondominant wrist. The device detects movement by means of a solid piezoelectric accelerometer with 0.35- to 7.5-Hz bandwidth and 0.5–2 G peak value. The device was set to record 30-second epochs at medium sensitivity to detect wake threshold. The actigraphy recording showed the regularity and timing of sleep periods, and the Actiware 6.0.7 software calculated all the sleep parameters, including sleep onset latency, wake after sleep onset, total sleep time and sleep efficiency.

The Consensus Sleep Diary was developed to provide a standard sleep log based on patients’ input and expert consensus. The sleep indices in the diary differentiate between people with insomnia and good sleepers. We used the diary for longitudinal subjective assessment of sleep pattern.

Biologic assessment of circadian phase

The dim-light melatonin onset test (Saliva Melatonin ELISA [enzyme-linked immunosorabent assay] kit [EK-DSM], Bühlmann Laboratories) comprises 8 saliva samples collected according to standard procedures. Three baseline samples were taken every 30 minutes, and the remaining samples were collected hourly. Data collection started 6 hours before habitual bedtime, which was determined based on participants’ self-report and was verified from their sleep diary data. Medications that influence melatonin levels were prohibited or discontinued before the test.

Saliva samples were frozen and analyzed by means of ELISA as per the manufacturer’s instructions. The intra- and inter-assay coefficients of variation were 7.62 and 8.88, respectively.
As per consensus guidelines, the time of dim-light melatonin onset was determined as the time at which the melatonin concentration reached and remained above the threshold (2 standard deviations above the baseline or, in cases in which there were fewer than 3 baseline values, an absolute value of 3 pg/mL).

### Diagnosis of circadian rhythm sleep–wake disorder

We diagnosed CRSWDs according to the third edition of the *International Classification of Sleep Disorders* based on agreement between a physician and a clinical psychologist (C.M.S. and D.M.Z.), both with specialization in sleep medicine. All 3 assessors were blind to the study hypothesis with the exception of the first author (D.M.Z.), who was involved in the administration of aspects of the assessment.

### Statistical analysis

Based on the rate of CRSWDs in mTBI of 36% observed by Ayalon and colleagues, an upper rate of CRSWDs generally associated with insomnia of 10%, 2-tailed $\alpha = 0.05$ and 95% power, we set 34 participants as a lower limit for recruitment to address our hypothesis.

We compared the observed proportion of CRSWDs with the population estimate of 10% using the $z$ test for 1 proportion, and 95% confidence intervals (CIs) for the prevalence rates were based on the corresponding precision afforded by our final sample size ($n = 50$).

### Ethics approval

This study was approved by the Ryerson University Research Ethics Board.

### Results

Participant selection is summarized in Figure 1. The final study group comprised 50 patients with mTBI and chronic insomnia symptoms. Participant characteristics and referral sources are summarized in Table 1.

Two participants had missing actigraphy data, and therefore we used their sleep diary data to determine whether they had a normal sleep phase. Three participants did not complete the dim-light melatonin onset test, and results could not be determined precisely from 5 samples (the onset was outside the measurement period in 3, and there were technical problems with 2 samples; these 2 were not included in the analysis).

Most participants (46 [92%]) had moderate or severe insomnia symptoms (Insomnia Severity Index score $\geq 15$). As reported during the interviews and verified by family when present, 37 participants (74%) did not have subjective sleep problems or sleep disorders before their injury.

Thirteen participants (26% [standard deviation 12%]) received a diagnosis of CRSWD, which supports our hypothesis of a greater proportion than the expected base rate of 10% for more general insomnia samples ($z = 3.77, p < 0.001$). The most common diagnosis was delayed sleep–wake phase disorder (10 patients [20%]). The mean time at dim-light melatonin onset for these patients was 2253 (95% CI 2218–2318), which represented a significant delay compared to those without a CRSWD (mean 2024, 95% CI 2006–2040, $t = 4.22, 32$ degrees of freedom, $r = 0.73$).

The median age of those with delayed sleep–wake phase disorder was 26 years (interquartile range [IQR] 20–30 yr); only 1 of these participants was older than 32 years. There were 2 participants with advanced sleep–wake phase disorder and 1 participant with irregular sleep–wake rhythm disorder; all 3 were more than 40 years. The median age of the 37 participants without CRSWDs was 40 years (IQR 31–54 yr).

There was no meaningful difference in the average Insomnia Severity Index score between participants with normally entrained circadian rhythm (19.8), those who had delayed sleep–wake phase disorder (20.7) and those who had advanced sleep–wake phase disorder (20.5). The Insomnia Severity Index score of the single participant with an irregular sleep–wake rhythm disorder was 24.

### Interpretation

We carried out a rigorous multimethod diagnostic circadian assessment based on standard criteria in a representative sample of patients with mTBI and chronic insomnia. The results support our hypothesis: 26% of the sample received a diagnosis of CRSWD. Previous circadian studies have included patients with moderate and severe TBI, assessed melatonin secretion without determining dim-light melatonin onset, or determined average melatonin onset in a group of patients without providing diagnostic assessment and interpretation, in samples that were substantially smaller than the current sample. The standard clinical intake assessment at least 3 months after the TBI ensured that a sample of people with chronic insomnia symptoms was included.
Ayalon and colleagues selected 15 participants from among 42 sleep clinic patients with mTBI and insomnia. A CRSWD was diagnosed in all 15 participants, which suggests that the prevalence of CRSWDs may be higher among patients with mTBI who have chronic insomnia symptoms than in relevant comparison groups (i.e., sleep clinic patients with insomnia). However, selection criteria, dim-light melatonin onset and time since injury were not specified, and the age range was restricted.

Most (77%) of the observed CRSWDs in our study reflected delayed sleep–wake phase disorder. The observed proportion of participants with delayed sleep–wake phase disorder, 20%, is 1–2 orders of magnitude higher than in the general population (0.17%–1.53%) and over 3 times that among patients with chronic insomnia treated at sleep clinics (6.7%). Estimates for adolescents are 3%–7% in the general population and 7%–16% in clinical samples of people with psychiatric conditions. The particularly high prevalence in the younger mTBI participants in the current study underscores the need for more studies to determine the rates of CRSWDs across demographic and clinical samples.

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This study did not explore the causes of delayed sleep–wake phase disorder, but several etiologic factors merit consideration. The authors of a magnetic resonance imaging study noted that patients who had sleep problems after an mTBI had a longer cerebellar tentorium and a flatter tentorial angle than patients who did not have sleep problems. They speculated that the pineal gland (the site of melatonin

| Characteristic                      | No. (%) of participants |
|------------------------------------|-------------------------|
| Age, yr                            | Median (IQR)            |
|                                    | 39.5 (23.8)             |
| Range                              | 17–62                   |
| Gender                             |                         |
| Male                               | 18 (36)                 |
| Female                             | 32 (64)                 |
| Education                          |                         |
| High school                        | 13 (26)                 |
| College                            | 12 (24)                 |
| University                         | 25 (50)                 |
| Employment status                  |                         |
| Full-time                          | 21 (42)                 |
| Part-time                          | 5 (10)                  |
| Student                            | 4 (8)                   |
| Unemployed                         | 8 (16)                  |
| Sick leave/disability              | 11 (22)                 |
| Retired                            | 1 (2)                   |
| No. of prior concussions           |                         |
| 0                                  | 32 (64)                 |
| ≥1                                 | 18 (36)                 |
| Time since injury, mo†             |                         |
| Median (IQR)                       | 11.5 (12)               |
| Range                              | 3–24                    |
| Cause of injury                    |                         |
| Car accident                       | 18 (36)                 |
| Sports injury                      | 18 (36)                 |
| Fall                               | 9 (18)                  |
| Hit on head by object              | 2 (4)                   |
| Physical assault                   | 2 (4)                   |
| Workplace injury                   | 1 (2)                   |
| Comorbidities‡                     | 22 (44)                 |
| Chronic pain                       | 13 (26)                 |
| Hypertension                       | 5 (10)                  |
| Gastrointestinal disease (e.g., gastrointestinal reflux) | 5 (10) |
| Hyperlipidemia                     | 3 (6)                   |
| Asthma                             | 3 (6)                   |
| Autoimmune disease                 | 2 (4)                   |
| Hypothyroidism§                    | 2 (4)                   |
| Osteoporosis                       | 1 (2)                   |
| Heart disease                      | 1 (2)                   |
| Past psychiatric condition         | 23 (46)                 |

| Characteristic                      | No. (%) of participants |
|------------------------------------|-------------------------|
| Current psychiatric condition      |                         |
| Major depressive disorder          | 13 (26)                 |
| Social anxiety disorder            | 7 (14)                  |
| Panic disorder                     | 4 (8)                   |
| Obsessive–compulsive disorder      | 2 (4)                   |
| Specific phobia                    | 1 (2)                   |
| Referral source                    |                         |
| Head injury clinic                 | 18 (36)                 |
| Family medicine clinic             | 16 (32)                 |
| Concussion/sports clinic           | 6 (12)                  |
| Sleep clinic                       | 4 (8)                   |
| Speech-language pathologist, neurologist, physiotherapist | 4 (8) |
| Emergency specialist               | 2 (4)                   |

Note: IQR = interquartile range. *Except where noted otherwise. †In participants with multiple concussions, time elapsed since the injury that preceded the onset of the sleep problem. ‡Some participants had multiple comorbidities. §Participants who reported a history of hypothyroidism were using thyroid supplements.
Research

Another biologic predisposing factor could be a susceptibility of the sleep and circadian system for phase delay owing to a long circadian period, heightened responsiveness to the melatonin-suppressing effect of light in the evening, or slow accumulation and dissipation of homeostatic sleep drive. If the social demand to wake up early in the morning lessens during the postinjury recovery period, the sleep phase follows the circadian signals and shifts to later hours in people with a predisposition for phase delay. This could lead to a vicious cycle in which longer light exposure in the evening and lack of light exposure in the morning further delay the circadian cycle.

It is possible that people with delayed dim-light melatonin onset or evening chronotype before their injury are at increased risk for a TBI because they are sleepy in the morning when they attend sport training, school or work. Athletic performance fluctuates across the circadian cycle, and excessive sleepiness affects cognitive performance and increases the risk for accidents and injuries. Thus, such susceptibility factors may partly contribute to a high prevalence of delayed sleep–wake phase disorder in mTBI.

Two participants in our study received a diagnosis of advanced sleep–wake phase disorder. Reports of this disorder are rare, including only individual cases and family cohorts. In the current study, neither of the participants with the disorder had a family history of this condition.

One participant was diagnosed with irregular sleep–wake rhythm disorder. This CRSWD has previously been described among people with brain disorders. The affected person in the present study had an alcohol use disorder until 6 months before the injury. There is an interaction between the circadian system and alcohol use, and it is possible that preinjury alcohol use disorder increases the risk of an irregular sleep–wake rhythm disorder following a brain injury.

Awareness of CRSWDs is generally low among clinicians, but distinguishing them from insomnia disorder is crucial since the treatment of insomnia disorder (cognitive behavioral therapy or selected hypnotic medications) and the management of CRSWDs (melatonin and bright light therapy) are fundamentally different.

Our results indicate that clinicians should be attentive to symptoms of CRSWDs when they see patients with persistent insomnia symptoms following an mTBI, including concussion. Referral for a circadian assessment is warranted if there is a marked shift of the sleep period or if the patient sleeps only for short (maximum 4-hr) periods around the clock. We suggest that the threshold for referring to a circadian assessment should be especially low if a teenager or young adult reports chronic sleep-onset insomnia or a delay of sleep phase following an mTBI.

The results of the current study highlight the need to assess CRSWDs in patients with chronic insomnia following an mTBI. Circadian assessment is currently not part of a standard investigation of postconcussion insomnia and is not a routine component of sleep clinic assessments. Consequently, based on our results, about 25% of patients with symptoms of postconcussion chronic insomnia would not receive an appropriate diagnosis and thereby tailored treatment for their sleep difficulty, which, in turn, would lead to lower likelihood of recovery.

Future longitudinal studies should identify factors that predispose people to a CRSWD after a concussion, and test the nature and direction of pathways between the circadian timing of sleep and wakefulness and factors that may influence this relation in mTBI in various age groups. Given the targeted selection process for this study, future research should aim to ascertain rates of CRSWDs through broader recruitment of mTBI populations. In addition, future studies should assess CRSWDs in larger samples of people with mTBI who have insomnia.

Replication of this study will benefit from inclusion of an age- and sex-matched comparison group of patients with chronic insomnia who did not have a history of TBI to allow for more direct comparison of rates of CRSWDs and sleep–wake measures. Given that research on CRSWDs among patients with insomnia alone is limited, future studies that include comparison groups will add valuable information not only for those with TBI but also to the general insomnia and CRSWD literature.

Of potential interest for future study, the median age of those with delayed sleep–wake phase disorder in the current study was 26 years (IQR 20–30 yr); only 1 of these people was older than 32 years.

Limitations

Some limitations of our study should be considered. First, the sample may overrepresent patients who are distressed by their insomnia symptoms. At the same time, these are the patients who (ideally) would receive some form of postconcussion sleep assessment in the community. Second, we did not recruit people with substance use disorders or shift workers. It is likely that the frequency of CRSWDs is higher in these groups than in our sample. Also, participants may not have reported insomnia symptoms that predated their injury. This does not bias the diagnoses but precludes inferences about the possible role of the injury. The results pertain to treatment-seeking people with chronic insomnia symptoms who had an mTBI 3–24 months before the sleep assessment and may not be generalizable beyond this group. Although sufficient for our current aims, the sample size limited precision in the estimated prevalence of CRSWDs. In addition, we did not include an age- and sex-matched comparison group of patients with chronic insomnia who did not have a history of TBI. The current study did not aim to establish a cause-and-effect relation between the clinical diagnoses and the injury, or control for variables that may have influenced this relation.

Conclusion

A CRSWD was diagnosed in about one-quarter of treatment-seeking people with symptoms of postconcussion chronic insomnia. This suggests that clinicians may include CRSWDs in their diagnostic algorithm and consider the potential benefits of a circadian assessment when patients report insomnia symptoms and a marked change in their sleep phase following an mTBI. The majority of participants in whom a CRSWD was diagnosed were adolescents and young adults with a...
delayed phase sleep–wake disorder; this finding warrants largescale replication. Given that insomnia symptoms are major predictors of poor postconcussion recovery, detection and effective management of CRSWDs may improve recovery in those who experience insomnia following an mTBI.

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Affiliations: Department of Psychology (Zalai, Girard), Ryerson University, Toronto, Ont.; Oskit Centre for Cognitive Therapy (Zalai), Oakville, Ont.; Division of Neurosurgery and Injury Prevention (Cusimano), St. Michael’s Hospital, University of Toronto; Department of Psychiatry (Shapiro), University of Toronto; Youthdale Child and Adolescent Sleep Centre (Shapiro), Toronto, Ont.

Contributors: Dora Zalai conceived the study. Dora Zalai and Colin Shapiro acquired the data, and Dora Zalai and Todd Girard analyzed the data. Dora Zalai drafted the manuscript. All of the authors contributed to the study design, data interpretation and revising the manuscript critically for important intellectual content, approved the final version to be published and agreed to be accountable for all aspects of the work.

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