Sleep disturbance in mental health problems
and neurodegenerative disease

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Abstract: Sleep has been described as being of the brain, by the brain, and for the brain. This fundamental neurobiological behavior is controlled by homeostatic and circadian (24-hour) processes and is vital for normal brain function. This review will outline the normal sleep–wake cycle, the changes that occur during aging, and the specific patterns of sleep disturbance that occur in association with both mental health disorders and neurodegenerative disorders. The role of primary sleep disorders such as insomnia, obstructive sleep apnea, and REM sleep behavior disorder as potential causes or risk factors for particular mental health or neurodegenerative problems will also be discussed.

Keywords: sleep, mental health, neurodegenerative disorders, cognition

Normal sleep physiology
Sleep is precisely defined by behavioral and electrophysiological measures. Electrophysiologically, sleep is distinguished from wake by distinct changes in the electroencephalogram (EEG), electrooculography, and muscle activity as measured by electromyography. Based on these measures, sleep has been divided into two states, rapid eye movement (REM) and non-REM (NREM) sleep. There are slow rolling movements on electrooculography during NREM sleep and rapid eye movements during REM sleep. REM sleep is characterized by higher-frequency EEG activity and an almost total loss of skeletal muscle tone seen on the electromyography.1 NREM sleep accounts for 75%–80% of total sleep time, predominates during the early stages of sleep, and is subdivided into three stages: N1 (drowsiness), N2 (light sleep), and N3 (deep sleep).2 The transition from wake to deep sleep through these stages is accompanied by a slowing of the EEG from high-frequency, low-voltage waves (beta waves) to low-frequency, higher-voltage (1–3 Hz) waves (delta waves), also called slow-wave activity, reflecting an increased cortical synchronicity.1 NREM sleep rapidly transitions into REM sleep 60–90 minutes or more after onset. During REM sleep, the EEG shows higher frequency lower amplitude waves in the theta range.1 There is regular transition between NREM and REM sleep in approximately 90–120 minute cycles with NREM sleep dominating during the early part of the night and REM later on. The timing and duration of sleep is controlled by two processes:3 the homeostatic sleep drive (process S), which increases for each hour of wakefulness, and the circadian rhythm (process C), where, naturally, humans sleep during the night and are awake during the day.

This sleep rhythm is driven by the central circadian pacemaker or master clock located in the suprachiasmatic nucleus (SCN) of the anterior hypothalamus.
At the cellular level, the master clock is controlled by an array of clock genes\(^4\) that ensure it runs at a period of just over 24 hours.\(^5\) The clocks are entrained to the 24-hour cycle by both environmental and endogenous cues called zeitgebers. Light is the major environmental zeitgeber facilitated by photosensitive retinal ganglion cells which have a direct neuronal link from the retina to the SCN via the retinohypothalamic tract.\(^6\)

**The role of sleep in mental health and cognition**

Important clues about the role of normal sleep in both memory function and mental health are derived from studies of sleep deprivation (SD) and sleep restriction (SR) in healthy subjects.

In healthy subjects, experimentally induced sleep loss is associated with impairments in a broad range of cognitive functions\(^7,8\) and people with primary sleep disorders such as insomnia\(^9\) and sleep apnea\(^10\) also suffer from cognitive impairments. Cognitive performance also follows a well characterized circadian rhythm,\(^11\) and people suffering from circadian rhythm disorders also suffer from impairments in cognitive performance.\(^12\)

Both SD, involving complete loss of sleep over several days, and SR, involving partial sleep loss, have been used in experimental studies to examine the effects of sleep loss on cognitive function. During studies of SD and SR, a multitude of cognitive tests have been performed examining a broad range of cognitive domains. Several meta analyses have been performed on the data from studies examining the effects of SD and cognition in healthy subjects.\(^8,13–15\) There are significantly fewer studies examining the effects of SR on cognition and no meta analyses performed to date.

To summarize the data from a large number of studies examining different aspects of attention, vigilance, and learning tasks under conditions of SD/SR, SD and SR consistently and significantly impair simple attention as evidenced by slowing of reaction time (RT) and increased lapses during simple vigilance tests. The effects of moderate SR accumulate substantially over time and can be equal to the effects of several days of total SD.

Performance within individuals demonstrates increasing variability over increasing time of SD, which is best explained by wake/sleep state instability and the interaction with circadian rhythm during sleep and wake. There is considerable variability between individuals with respect to their susceptibility to the effects of SD and SR, although the mechanisms are as yet unknown. Executive functions are less susceptible to the effects of SD and SR than simple attention and deficits are found less consistently in studies. Logic and rule-based tasks measuring crystallized abilities are not substantially affected by SD. In contrast, tasks involving more divergent and innovative thinking are significantly impaired by SD.

Sleep appears to be necessary for both memory encoding and consolidation and therefore learning and memory are significantly affected by SD both before and after learning. Emotional memory is also susceptible to the effects of SD, which may lead to relative preservation of negative compared to positive and neutral memories.

**Sleep disturbance in mental health problems**

**Primary sleep disorders as causal factors in mental health disorders**

Insomnia is the most common primary sleep disorder and is defined as difficulty falling asleep, staying asleep, or nonrestorative sleep despite adequate opportunity to sleep. Within population-based studies, these symptoms alone affect approximately 30% of adults.\(^16\) When one adds associated daytime impairment or distress as a function of the insomnia and symptoms lasting over 1 month that are not secondary to another sleep disorder or mental disorder, then prevalence estimates range from 6%–10%.\(^17–20\) The place of insomnia within both the International Classification for Sleep Disorders and the *Diagnostic and Statistical Manual of Mental Disorders* highlights the interface between disturbed sleep and the psychological distress associated with that sleep disturbance.\(^21,22\) Primary insomnia, unrelated to another medical or psychiatric condition, is considered to be part of a psychophysiological hyperarousal process. Age and sex are the best established risk factors, with increased prevalence in women and older adults. However, comorbid medical conditions,\(^23\) psychiatric illness,\(^24\) and working night shifts or rotating shifts all represent independent risk factors for insomnia. These conditions are thought to be insomnia precipitants in those already predisposed to developing sleep disturbance. The most common comorbidities associated with insomnia are psychiatric disorders, and it is estimated that 40% of all insomnia sufferers have a coexisting psychiatric condition.\(^24,25\) Among these psychiatric disorders, depression is most common, and insomnia is a diagnostic symptom for depressive and anxiety disorders.\(^26\) Insomnia is increasingly recognized as an independent risk factor for both dysthymia and major depressive disorder. A number of studies, including longitudinal studies in the young and
and a subsequent meta-analysis, have shown that those with insomnia are at least twice as likely to develop depression compared to people with no sleep difficulty over the subsequent 1–3 years of follow-up. Those with insomnia comorbid with depression are more likely to remain depressed despite standard treatments; therefore, depression and anxiety may be consequences of as well as risk factors for disrupted sleep. In particular, the hypervigilance and increased arousal associated with both anxiety and insomnia often lead to bidirectionally intertwined disorders that interfere with both sleep onset and sleep maintenance. This highlights a potentially underused treatment for depression. There is robust evidence base for cognitive behavioral therapy for insomnia, with many prospective controlled trials showing sustained benefit whether insomnia is primary or comorbid with other medical or psychiatric conditions. Importantly, a number of these studies in patients with associated depression and anxiety show an effect regardless of baseline depression levels and moderate treatment effects on both anxiety and depression outcomes as well as insomnia scores.

**Obstructive sleep apnea (OSA)**

Snoring and associated pauses in breathing (but not snoring alone) was strongly associated with major depression in a large US population study. Depression and neurocognitive symptoms have been associated with OSA but the pathophysiology remains poorly understood and some but not all studies identify OSA as independently associated with depression when other variables are controlled for. Many of the symptoms of OSA, such as fatigue, low energy, daytime sleepiness, poor concentration, and neurocognitive impairment, overlap with those of depression and may lead to diagnostic difficulty. Improvement of many of these symptoms, including depression measures and quality of life, can be demonstrated with continuous positive airway pressure treatment, making it important to consider OSA as a potential diagnosis or modifiable factor in those with mental health problems.

**Schizophrenia**

Disturbed sleep is found in 30%–80% of patients with schizophrenia, depending on the degree of psychotic symptomatology. Sleep disturbance is inversely correlated with quality of life, and a large number of polysomnographic studies have shown increased sleep latency, reduced sleep consolidation, reduced total sleep time, and increased wake time after sleep onset. The sleep disturbances appear to be an important part of the pathophysiology of schizophrenia. These effects have been consistently reported in drug-naive patients, those who have withdrawn from medication, and those being treated with antipsychotics and other psychotropic medication.

Analysis of EEG and in particular dream sleep in schizophrenia patients has been of interest to psychiatrists as far back as 1955, when Dement first described a reduction in dream recall in patients with schizophrenia and abnormal REM parameters. Further studies have shown reduced REM sleep latency, higher REM densities, and a failure of REM rebound after sleep deprivation. However, there are several contradictory results in these studies, and REM sleep variables are not consistently affected.

A number of sleep parameters, such as the amount of slow-wave sleep (SWS) and REM latency, are significantly correlated with clinical variables, including severity of illness, positive symptoms, negative symptoms, outcome, neurocognitive impairment, and brain structure. Reduced SWS has been reported in patients with schizophrenia in a number of studies; however, there have been contradictory results in patients who were off medication or had never been treated. More detailed analyses of the spectral composition of the EEG have revealed a consistent and significant reduction of the higher-amplitude and lower-frequency delta waves, especially in the anterior frontal areas. Significant negative correlations have been reported between reduced SWS or high-amplitude, low-frequency delta waves and both negative symptoms and neurocognitive impairment. Given the need for SWS in the consolidation of memory, this offers a potential cause or contributory factor for the neurocognitive impairment in schizophrenia.

In schizophrenia patients, memory consolidation has been shown to be impaired compared to controls and to be positively correlated with the amount of SWS and sleep efficiency. Manoach et al measured the effects of sleep on procedural memory consolidation in schizophrenia patients. Daytime practice improved performance on a finger tapping motor sequence task equally in schizophrenia patients and controls whereas an improvement following overnight sleep was only found in the control group. In a subsequent study, consolidation in schizophrenia patients was dependent on both SWS and stage two sleep in the last quarter of the night. Therefore, the sleep abnormalities present in schizophrenia may interfere with normal sleep-dependent memory consolidation and therefore have an influence on cognitive function.

The direct effect of antipsychotic medication is unclear, and the confounding effect of neuroleptics is a major problem...
with many studies. First-generation or typical antipsychotics are associated with increased total sleep time, increased sleep continuity, and increased REM latency but the reported effects on other sleep stages are variable. Treatment withdrawal is followed by a change in sleep structure, mainly in the opposite direction, with a deterioration of sleep quality. There are no consistent effects of first-generation antipsychotics on measures of sleep continuity and sleep structure, including the percentage of sleep stages or sleep and REM latency in healthy controls. Therefore, it is possible any effects of these high-potency typical antipsychotics may be indirect with improved sleep secondary to reduced stressful symptomatology.

In contrast, studies of the effects of the atypical antipsychotics (clozapine, olanzapine, quetiapine, risperidone, ziprasidone and paliperidone) in healthy controls and schizophrenia patients show a relatively consistent effect on measures of sleep continuity, with an increased total sleep time and/or sleep efficiency and individually varying effects on other sleep parameters, such as an increase in REM latency observed for olanzapine, quetiapine, and ziprasidone and an increase in SWS documented for olanzapine and ziprasidone. Additionally, clozapine and olanzapine demonstrate comparable influences on other sleep variables, such as SWS or REM density, in controls and schizophrenic patients. Therefore, it is possible that the effects of second generation antipsychotics observed on sleep in healthy subjects and schizophrenic patients might involve the action of these drugs on symptomatology, such as depression, cognitive impairment, and negative and positive symptoms.

Circadian rhythm abnormalities have been described in a number of small studies using actigraphy, often without matched controls. Mills et al showed robust circadian rhythms of both core body temperature and urinary electrolytes but a shorter than 24-hour sleep–wake cycle. Wirz-Justice et al demonstrated normal rest activity cycles in clozapine-treated patients compared to disordered circadian rhythms in those on typical neuroleptics, and proposed that mechanisms of drug action directly affected circadian rhythm. Decreased physical activity during the day was also shown using actigraphy in schizophrenia patients compared to both controls and depressive patients. Bromundt et al showed highly variable sleep–wake cycles in 14 patients with more robust sleep–wake cycles correlating with better frontal lobe function, again highlighting a link between impaired sleep and worse cognition. The most recent and detailed analysis of circadian rhythm, which included prolonged actigraphy and melatonin profiles in 20 schizophrenia patients compared to 21 unemployed controls, found marked circadian rhythm abnormalities, including both delayed sleep-phase syndrome and free-running, non-24-hour circadian rhythm in 50% of those studied, with no relationship to the neuroleptic used.

A very recent study demonstrating disrupted sleep–wake rhythms in a mouse model of schizophrenia demonstrated intact retinal inputs to the SCN, but suggested a possible impairment of synaptic connectivity modulating the output from the SCN. Mouse models offer the potential for targeting the sleep and circadian rhythm disruption as a therapy for schizophrenia.

Disrupted sleep and wake in schizophrenia may be intrinsic to the disease itself or occur as a consequence of psychotropic medication, but it may also be due to a primary sleep disorder. This remains a somewhat neglected and potentially treatable aspect of sleep disturbance within this population. There is now increasing recognition of the association between schizophrenia and sleep-related breathing disorders, possibly related, in part, to the weight gain associated with atypical antipsychotics. Metabolic syndrome was found in 38% of a community-based psychiatric cohort on long-term antipsychotics. A number of studies have found high rates of sleep apnea in both hospital and community-based populations. However, not all studies have used control groups, and it has been suggested that sleep apnea is so common in the increasingly obese general population that these rates may not differ to those of BMI matched controls. To the authors’ knowledge, there have been no cross-sectional or prospective studies of psychiatric patients with metabolic syndrome to assess the prevalence of OSA. Previous inpatient studies have shown that obese patients with chronic psychiatric morbidity have high rates of OSA. Occasional case reports have highlighted the benefits of continuous positive airway-pressure treatment on symptomatology in schizophrenia but there are no prospective trials to assess long-term benefit or tolerability of continuous positive airway-pressure therapy in comparison to normal controls.

There are some hypotheses that periodic limb movements are not increased in those with schizophrenia, but there is very little data regarding the prevalence of other primary sleep disorders, including restless legs and parasomnias.

With regards to insomnia, symptoms are common in those with high degrees of paranoia and, recently, the first structured pilot trial of cognitive behavioral therapy for insomnia was carried out in 15 patients with fixed and treatment-resistant psychosis. This showed a promising outcome with a significant improvement not just in sleep outcomes in two-thirds of subjects, but reduced persecutory
delusions in 50%. Larger and longer-term trials are needed with control groups to assess the long-term benefits of such behavioral therapies.

**Major depressive disorder**

Sleep complaints and depression are bidirectionally related, and there has possibly been more general focus on the role of sleep disturbance in depression than in any other psychiatric disorder with sleep disturbance included within the *Diagnostic and Statistical Manual of Mental Disorders Fourth Edition* (DSM-IV) diagnostic criteria. Epidemiologic studies confirm poor sleep quality in 50%-90% of those with diagnosed depression, and the high prevalence of depression in insomnia patients has already been discussed, with insomnia emerging as a clear risk factor for subsequent depression and conferring a worse outcome in those already depressed. Despite methodological differences, sleep disturbance is one of the most important predictors of a subsequent depressive relapse. The discovery that complete SD exerts a potent antidepressant effect therefore seems paradoxical, but much of the original interest in the relationship between sleep and mood stemmed from experiments in the 1970s by Wirz-Justice and Van den Hoofdakker, among others, to show that a single night of complete SD leads to a rapid and dramatic improvement of mood in up to 60% of patients. However, the effect is short-lived, with immediate relapse after normal sleep, sometimes after even a brief nap. The polysomnography changes that occur during depression have been studied from the 1960s, and a number of studies have consistently shown a reduction in sleep efficiency, decreased SWS and disinhibited REM sleep with shortened REM latency, a longer first REM cycle, and increased REM density (the number of eye movements during REM). Decreased REM latency was initially thought to be a marker for primary rather than secondary depression, but subsequent studies failed to show the same result. More detailed analysis of SWS has shown a reduction in delta waves, particularly in the first half of the night and correlated to symptom severity. However, to date, no single sleep variable has been of sufficient specificity to be diagnostic for any one psychiatric illness.

One key question has been whether sleep abnormalities only occur acutely during depressive relapse as a “state-dependent” marker, or are present prior to symptom onset and confer a vulnerability to subsequent mood disorder as a “trait” marker. However, this area is complicated by the effects of medication on polysomnography studies. Researchers have shown REM sleep normalizes after depression treatment, but many of the pharmacotherapies and in particular the antidepressants are known to lengthen REM latency and improve sleep continuity in both patients and normal controls. However, increased sleep efficiency and REM sleep improvement has also been demonstrated after nonpharmacological therapies such as cognitive behavioral therapy. Conversely, several studies have shown that changes such as shortened REM latencies persist during remission, and longitudinal studies have shown a stable REM latency over time, suggesting at least some trait markers of at least some of the sleep changes. In further support of a trait marker is the finding of similar REM sleep changes in subjects with a strong family history of depression but no symptoms at the time of study. Although up to 20% of patients complain of hypersomnia, there are few polysomnographic studies of this group, and multiple sleep latency tests and measured total sleep time in this group are often normal. The same is true of seasonal affective disorder, despite complaints of hypersomnia, suggesting that this is not the same sleepiness that can be objectively demonstrated in conditions such as idiopathic hypersomnia. Possible confounding factors include different subtypes of depression within studies and variable tools used to assess daytime sleepiness.

**Effect of antidepressant therapies on sleep**

Total SD has been mentioned earlier as the only therapy with proven effect within 24 hours; however, rapid relapse is the norm after a single night of recovery sleep, making it of little therapeutic benefit. Both timing and dose of SD have been used to try and obtain therapeutic benefit, with little success to date. Selective REM suppression has been used, and one early study showed a treatment response comparable to imipramine, but later studies still showed greater benefit with total SD. Bright light may add some further benefit in depression, but there is better evidence for its benefit in seasonal affective disorder. Almost all antidepressant medications modify sleep architecture, making sleep changes in patients in remission hard to interpret. Most tricyclic antidepressants increase total sleep time and decrease wake time after sleep onset, while many selective serotonin reuptake inhibitors have the opposite effect. However, almost all antidepressants prolong REM sleep latency and reduce the amount of REM sleep. Both depressed patients and healthy volunteers have a decreased dream-recall frequency under treatment with antidepressants. This is a consistent effect in tricyclic antidepressants, less
consistently documented for selective serotonin reuptake inhibitors/serotonin norepinephrine reuptake inhibitors. Withdrawal from antidepressants may cause nightmares and intensify dreaming; however, this antidepressant effect is not dependent upon REM sleep elimination and there is no clear evidence that the mechanism of action of antidepressants is dependent on REM sleep suppression.

The role of circadian rhythm abnormalities in the pathophysiology remains unclear. While elevated cortisol is consistently seen in at least 50% of depressed patients at night, levels fall in remission and the effect is not specific to depression. Some advocate regular sleep–wake rhythms and avoidance of SD to avoid relapse. One recent study showed that a chronotherapeutic regime (bright light and sleep stabilization) significantly improved outcomes over a 9-week period but there is a lack of longitudinal data. It is clearly an area that warrants further study with well-defined diagnostic groups.

Bipolar disorder
Sleep disturbance is a core feature of bipolar disorder (BD), being included in DSM-IV diagnostic criteria for major depressive episodes, mania, and hypomania, and it is also commonly reported between mood episodes. Disturbances in circadian rhythm are also present, and it is now thought that sleep and circadian disturbances, rather than being an artifact of mood episodes, are a core part of the underlying etiology and maintenance of BD. This view is supported by the evidence of sleep and circadian disturbances in the initial prodrome of the illness and in the prodrome of relapse. Children at high risk for developing BD have increased sleep disturbances preceding the first episode, including difficulty getting to sleep, fragmented sleep, and decreased sleep. Sleep disturbances are reported as an early symptom of bipolar depression (17%–57% of subjects [median 24%]) and are the most robust prodromal symptom of mania reported by 53%–90% (median 77%) of subjects preceding a manic episode. Enforced SD can lead to the onset of manic or hypomanic episodes, although the literature is inconsistent, probably due to poor characterization of unipolar and bipolar subjects and post hoc follow-up. However, in a well-defined group of 206 inpatients with DSM-IV- and DSM-IV-defined bipolar depression, switch rates of 4.85% to mania and 5.83% to hypomania were observed following three cycles of total SD.

Sleep during mania
Polysomnography studies consistently found shortened sleep time, shorter REM latency, increased REM density, and more disturbed sleep. Manic patients report a decreased need for sleep in 69%–100% of subjects studied.

Sleep and bipolar depression
Sleep disturbances during bipolar depression are common but follow a less consistent pattern than during mania. Hypersomnia is reported in between 23% and 78% of subjects and rates of insomnia vary considerably, with one study of patients. Polysomnography studies comparing patients with bipolar depression to unipolar depression and healthy controls report inconsistent results with greater fragmentation of REM sleep, lower REM latency, no difference in REM latency, increased REM density, fewer minutes in stage 1 sleep, and prolonged SWS latency all reported. The decrease in REM latency and higher REM density compared to healthy controls is probably the most consistent finding.

Sleep during euthymia
Polysomnography studies show greater REM density in euthymic BD patients than in controls, as well as more overnight arousals with increased REM sleep. Clinically significant sleep disturbance was found in 70% of
euthymic BD patients with 55% meeting diagnostic criteria for insomnia by Harvey et al."112 Compared to insomniacs and good sleepers, BD patients had lower sleep efficiency and lower daytime activity levels. Subjectively, BD patients report sleep difficulties that are more severe than in controls but less severe than in the insomnia group, and suffer more daytime sleepiness according to the Epworth Sleepiness Scale.137 Hypersomnia has also been reported in euthymic BD patients with 25% of patients meeting the criteria in one study.138

In summary, studies of euthymic BD patients indicate that there are significant sleep disturbances and variability in sleep, present in the absence of major mood episodes.

Sleep function and course of illness in BD

Given the possibility that sleep disturbances may have a causal role in the relapse and maintenance of bipolar disorder, several studies have examined associations between sleep function and course of illness. In a polysomnography study, REM and stage 2 sleep were not found to correlate with current symptoms, but duration of the first REM period and amount of SWS did correlate positively with manic symptoms and impairment measured on the Work And Social Adjustment Scale 3 months later.139 REM density was positively correlated with depressive symptoms and impairment at 3 months, and the amount of stage 2 sleep was negatively correlated with manic symptoms and impairment. In a study utilizing sleep diaries and semi-structured interviews,135 lower and more variable sleep efficiency and more variable total wake time were associated with more lifetime depressive episodes. Variability in time to fall asleep was positively correlated with concurrent manic symptoms. Hypersomnia has also been found to correlate with future depressive symptoms at 6 months.138 These studies suggest that sleep architecture may have a mechanistic role in disease process.

Circadian rhythms in BD

Circadian rhythm disturbance is reported in BD and may be at least partially responsible for the pattern of disturbed sleep evident in BD patients. Several actigraphy studies have found evidence of disturbed circadian rhythms in BD patients, including phase advances and greater variability in sleep patterns.140,141 A number of circadian genes have been linked to susceptibility to developing BD, including CLOCK, ARNTL 1-2, PER1-3, and CRY1-2.142 Overall, however, findings associating circadian genes to BD are inconsistent. Several reports indicate BD patients are more likely to be evening types,143-146 suggesting a circadian phase delay in BD. However, actigraphy-based studies have found variable results, with advanced sleep phase, delayed sleep phase, and very irregular sleep–wake cycles all found, as well as some groups showing few abnormalities. One consistent finding was reduced total activity compared to controls.141,147 Poor social rhythm regularity predicted the time to the next mood episode.148

Differences in the secretion of melatonin, and sensitivity of melatonin secretion to light, have also been reported in BD. Earlier studies149-151 suggested a supersensitivity to light in BD patients, with nighttime melatonin levels falling more compared to controls; later, larger studies did not find these changes, but did consistently show lower peak levels of melatonin with and without light stimulus and a less robust circadian rhythm to melatonin secretion.152-154 Recent evidence of lower activity of a key enzyme involved in melatonin synthesis (acetyleserotonin O-methyltransferase) in BD patients compared to controls suggests a general role
of melatonin as a susceptibility factor for the development of BD.\textsuperscript{155} Ramelteon, a selective melatonin MT1/MT2 agonist, may be effective in preventing relapse in euthymic bipolar patients,\textsuperscript{156} and agomelatine may be effective as an additional therapy for the treatment of bipolar depression.\textsuperscript{157} Collectively, these studies demonstrate abnormalities in circadian rhythms in patients with BD. Therefore, chronotherapeutics (controlled exposure to environmental stimuli that act on biological rhythms), such as SD, light, and dark therapy may be useful interventions for BD and some small studies have shown this to be the case.\textsuperscript{113,158,159} Studies have also demonstrated the efficacy of interpersonal social rhythm therapy, which aims to stabilize social rhythms in preventing BD relapse.\textsuperscript{160,161}

**Primary sleep disorders in BD**

The prevalence of sleep apnea in BD patients has not been systematically evaluated, despite the high frequency of risk factors such as overweight and obesity.\textsuperscript{162,163} A recent study using a self-assessment tool to establish the risk of OSA found 54.1\% of 72 bipolar I disorder patients were at a high risk of OSA. Despite a number of pharmacotherapies for BD being known to exacerbate restless legs, there have been very few studies of the prevalence of restless legs or periodic limb movements.

**Neurodegenerative diseases**

**Aging and sleep**

Given the strong association of neurodegenerative conditions with aging, it is important to understand the changes that occur within the sleep–wake cycle and circadian rhythm in an older population. As we age, total sleep time is slightly reduced compared to younger adults, with decreasing amounts of SWS, increased sleep latency, and increased sleep fragmentation during the night.\textsuperscript{164}

Disruptions in the sleep–wake cycle and sleep complaints are commonly found in community-based studies of older people. Over 50\% of adults aged over 65 have at least one chronic sleep-related problem.\textsuperscript{165,166} Older adults also have high rates of primary sleep disorders. Sleep-related breathing disorders, in particular OSA, are estimated at rates of 10\%–20\%.\textsuperscript{167–170} Insomnia and restless legs are also increasingly frequent in the people over 65.\textsuperscript{171,172} Changes in circadian rhythms have also been demonstrated with advancing age, with a decline in the cortisol and melatonin rhythms that entrain day–night activity patterns.\textsuperscript{173,174} With aging there is reduced amplitude of the circadian rhythm, a phase shift (in particular a phase advance),\textsuperscript{175} and a loss of the robustness of the rhythm (weakening of the rhythmic pattern). However, within older populations, there is significant circadian rhythm variability between individuals, with some hypothesizing that an age-related disruption to the SCN is responsible.\textsuperscript{176}

**Alzheimer’s disease (AD)**

Sleep disturbance is one of the most common reasons for institutionalization of demented elderly patients.\textsuperscript{177} From 25\%–50\% of the demented elderly suffer from severe nocturnal restlessness, often called “sundowning,” at some stage of the disease. Activity monitoring (actigraphy) in both early and late AD\textsuperscript{178–180} and in other dementias in patients at home and in institutions confirms profoundly disrupted circadian rhythms in those with moderate and severe dementia.

A recent large trial of bright light therapy and melatonin in institutionalized patients showed modest benefits in measures of daytime function with light therapy alone and together with melatonin;\textsuperscript{181} however, a further trial of melatonin alone showed no significant improvement in daytime function or night sleep.\textsuperscript{182}

In patients with AD, certain changes can be seen as an exaggeration of the normal aging changes with polysomnography showing increased numbers of awakenings and therefore increased stage 1 sleep. Compared to normal controls there is also reduced SWS.\textsuperscript{183–186} These changes are consistently reported across moderate to severe AD patients and sleep disturbance worsens with increasing severity of AD. The apolipoprotein status has been associated with progression of sleep–wake disturbance.\textsuperscript{187} Another feature of the polysomnogram in AD that suggests accelerated aging is a loss of some of the distinctive features of stage 2 sleep with poorly formed sleep spindles that are of shorter duration and less numerous.\textsuperscript{188} As the disease progresses, NREM sleep stages become progressively more difficult to stage, although REM sleep duration remains relatively stable and loss of REM atonia or significant REM sleep behavior disorder is uncommon.\textsuperscript{189}

**Parkinson’s disease (PD)**

In his elegant monograph published in 1817, James Parkinson gave his name to a progressive neurological disease characterized by stiffness, slowness, and tremor.\textsuperscript{190} In the last 10 years, there has been increasing clinical and research interest in the nonmotor symptoms of the disease. Sleep disturbance, depression, and cognitive impairment are all frequently seen in PD and are often considered to be more disabling than the abnormal movement.\textsuperscript{191,192} Sleep disturbance is near universal in advanced disease and some sleep symptoms can predat
all other motor phenomena. Particular mention should be made of the association between REM behavior disorder (REMBD) and Parkinsonian syndromes.

Carlos Schenck first described a group of patients with apparently idiopathic REMBD who went on to develop Parkinsonian syndromes including dementia with Lewy bodies (DLB), PD, and multiple system atrophy.

REMBD affects approximately 0.5% of elderly males, with patients acting out increasingly violent with vivid dreams that often result in injury to themselves or their bed partners. It is frequently well treated with long-acting hypnotics such as clonazepam. It is now known that a patient presenting to the sleep clinic with typical REMBD will have a 50% chance of developing a Parkinsonian syndrome within 5 years. A number of other symptoms can predate the onset of PD such as impaired olfaction, constipation, and mood disturbance but REMBD has the highest specificity as a biomarker raising the possibility of targeting coming neuroprotective agents at patients with this condition. Some patients, however, can develop symptoms many years before the development of any other neurodegenerative problem. Conversely, 40% of patients in a movement disorders clinic with PD will have REMBD and 95% of those will have multiple system atrophy. Interestingly, this association between REMBD and subsequent neurodegeneration is much more strongly associated with alpha synucleinopathies, including PD, multiple system atrophy, and DLB, but not with tauopathies, such as AD or progressive supranuclear palsy where a far smaller percentage of patients have loss of REM atonia or symptomatic REMBD and this diagnosis has been shown to improve differential diagnosis with these conditions. There is an increasingly clear association between REMBD and increased risk of the subsequent development of dementia and daytime visual hallucinations.

Daytime sleepiness was also highlighted in PD patients, particularly with the advent of newer dopamine agonists and the growing reports of “sleep attacks” in association with this group of drugs. Patients with PD need to be cautioned about possible daytime sleepiness when they start dopamine agonists. However, daytime sleepiness in PD is often multifactorial with many case control studies highlighting a large number of potential causes including severe REMBD, nocturia, and nighttime pain caused by immobility and tremor. Increased rates of both central and OSA have been described in those with moderate and advanced PD and there is also an increase in periodic limb movements and restless legs. Although some authors have suggested a possible disruption of the circadian rhythm in those with PD, to date, controlled studies are lacking. There is also a relative lack of prospective studies to determine which sleep disorders emerge at different stages of the disease and which therapies best improve night sleep and subsequent daytime function.

A careful sleep history should be taken in all patients with PD and sleep symptoms monitored throughout the disease. PD must be reported to the Driver and Vehicle

Figure 3 A 30-second epoch from a polysomnograph demonstrating the changes in REM behavior disorder.

Notes: The electrooculogram channels (LOC and ROC; blue) show the characteristic eye flicks of REM sleep. The submental electromyogram (CHIN 1; red) should be flat but shows phasic activity as the patient is moving.

Abbreviations: LOC, left outer canthus of the eye; REM, rapid eye movement; ROC, right outer canthus of the eye.
Licensing Authority in the UK and clinicians must include an assessment of sleepiness. Recent data show that PD patients themselves are poor judges of their safety behind the wheel but a recent review highlights the lack of standardized criteria to ensure safety behind the wheel.

Dementia with Lewy bodies (DLB)

There is less information on nocturnal sleep disturbance in DLB compared to AD, but recent studies suggest an even greater level of nighttime sleep disturbance with an increase in hallucinations, agitation, and apathy but a less clear relationship to disease progression. RBD remains a distinguishing feature within the history of a dementing patient that allows for greater sensitivity in diagnosis when present, but there is very limited data on other primary sleep disorders within this group alone. A recent large cross-sectional study across a number of dementia subtypes highlights RBD as distinguishing DLB and PD dementia from vascular dementia and AD and also the need to look for other sleep disorders, such as sleep apnea, which were frequent, particularly in vascular dementia.

Frontotemporal dementia

A single study using actigraphy and sleep diaries showed disturbed circadian rhythms in frontotemporal dementia, but this was less marked than that seen in AD and did not clearly relate to disease progression. Comparison between the polysomnography studies of those with AD compared to frontotemporal dementia showed greater REM sleep disruption in AD, again suggesting that sleep in frontotemporal dementia may be better preserved.

Huntington’s disease (HD)

HD is a well-described hereditary, neurodegenerative disease characterized by cognitive decline, behavioral change, and chorea, and patients often have particularly fragmented sleep, with up to 88% reporting sleep problems. A wide range of sleep problems, including increased movements during sleep, prolonged sleep latency, and nocturnal waking and daytime sleepiness, have all been reported. Unlike a number of other neurodegenerative conditions, patients with HD showed increased density of sleep spindles compared to healthy controls. The sleep disturbance correlates with the degree of cognitive impairment and depression.

Studies looking at the frequency of sleep-disordered breathing have shown variable results. A small pathophysiological study of five patients showed a subtle reduction of hypocretinergic neurons (27%) postmortem but cerebrospinal fluid hypocretin levels were normal in two studies of HD patients. Polysonomography studies of HD patients have shown reduced REM sleep and delayed REM sleep latency in those with established disease but also in premanifest disease with an increase in periodic limb movements, sleep apnea, and daytime sleepiness.

Summary

Sleep is vital for normal brain function and there is a complex bidirectional relationship between disturbed sleep and cognitive and mental health disturbance. Much research still needs to be done to characterize the different sleep patterns that occur within different diseases to see whether sleep patterns can be used as specific biomarkers of disease or, possibly more importantly, as trait markers for subsequent illness. It is still unknown whether intervening to normalize sleep clearly improves the outcome of chronic mental health problems or neurodegenerative disease, and there is an urgent need for increasing recognition of the role of disturbed sleep in a disturbed brain.

Disclosure

The authors report no conflicts of interest in this work.

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