Sleep, Circadian Rhythms, and Alzheimer’s Disease

Sleep and circadian rhythm disturbances are symptoms of Alzheimer’s disease that may also contribute to disease pathogenesis.

By Erik S. Musiek, MD, PhD

The high prevalence and public impact of Alzheimer’s disease (AD) make research efforts to uncover potential modifiable risk factors critically important. Sleep and circadian rhythm disturbances are common in patients with AD, especially as the disease progresses. However, an emerging literature supports the idea that sleep and circadian rhythm disturbances occur very early in the disease and may influence AD pathogenesis. This suggests that disrupted sleep and circadian timing may be modifiable risk factors. Data from mouse models show a strong relationship between sleep and the regulation of amyloid-β peptide (Aβ), the principal component of the amyloid plaques that define AD pathologically. In humans, acute sleep deprivation can drive increased Aβ levels, further supporting this relationship. Chronic sleep problems are associated with AD neuropathology, although a causal relationship has not been definitely proven.

Sleep and Circadian Rhythms in AD Dementia

Sleep quality and timing tend to change as individuals age and can be severely disrupted in AD dementia (ADD). The timing of sleep is strongly influenced by the circadian system, which generates 24-hour rhythms in many biological processes, synchronizing these to the external light-dark cycle. People with robust circadian timing enjoy consistent waketimes and bedtimes, active days, and restful nights. Although sleep is separate from the circadian clock, the circadian system prompts sleep at night by timing release of melatonin and direct signaling to sleep nuclei in the brain. Disrupted circadian function can lead to fragmented sleep timing, often manifested as erratic sleep at night and frequent napping during the day. This pattern is often observed in patients with ADD and can range from mild fragmentation of sleep timing to total breakdown of daytime boundaries. Patients with AD have neuronal degeneration in the suprachiasmatic nucleus of the hypothalamus, the master circadian clock of the body, that may underlie this fragmentation. Patients with AD also have changes in sleep itself, including decreases in both rapid eye movement (REM) sleep and slow-wave nonREM sleep. Changes in circadian sleep timing and sleep quality can be a major problem for caregivers and are a leading cause of institutionalization for patients with AD.

Sleep and Circadian Rhythm Changes in Preclinical Alzheimer’s Disease

Although sleep and circadian disturbances in ADD have been long-appreciated, it is only recent advances in biomarkers of AD that allow researchers to investigate these changes very early in the disease course, even prior to the onset of clinical symptoms. Longitudinal biomarker studies show that amyloid plaque pathology is present for many years, perhaps even decades, before cognitive symptoms occur. Increased levels of tau protein in the cerebrospinal fluid (CSF), another pathologic hallmark of AD, also precede the onset of cognitive decline by a few years. Individuals with evidence of amyloid or amyloid and tau pathology are now defined as having preclinical AD in research studies. Preclinical AD is associated with self-reports of poor sleep, difficulty going to sleep, and excessive daytime sleepiness, and objective measures of poor sleep efficiency (ie, more time in bed without sleep) and increased daytime napping. Fragmentation of circadian rhythms during activity is also detectable in preclinical AD, suggesting that AD-related changes to the circadian system may underlie some of the observed changes in sleep. Taken together, these data make it clear that sleep and circadian disturbances begin during preclinical AD and may predict eventual ADD, although longitudinal studies are still needed to define causality of such a relationship.

Sleep, Circadian Changes, and Risk of Alzheimer’s Disease

Epidemiologic researchers have examined whether sleep quality or duration over a lifetime influences risk for
having AD. Studies in adults over age 65 associate poor sleep with increased risk of developing dementia, although it is unclear if this reflects undiagnosed preclinical AD, or if sleep changes precede amyloid plaque deposition. Long average sleep duration (ie, > 12 h/night) is also associated with increased risk of developing symptomatic AD. However, this effect is seen primarily in individuals who likely had early-stage AD already. One interpretation of this correlation is that people begin to sleep longer in the early stages of the disease rather than long sleep being a risk factor earlier in life. Fragmented sleep and circadian rhythm function can precede the onset of symptomatic dementia. Sleep fragmentation and less robust circadian function in adults with an average age of 81 to 82 years who had no cognitive impairments are both associated with increased risk of developing AD within 5 years. Changes in sleep and circadian function, in particular daytime sleepiness, fragmented sleep at night, and increased daytime sleep, may be indicators of impending ADD.

Mechanisms Linking Sleep and Circadian Rhythms to Alzheimer’s Disease

Mouse models of AD are providing insights into how sleep and circadian disruption may play a causative role in the development of AD. As a result of synaptic activity, neurons release Aβ and aggregates in a concentration-dependent manner to form amyloid plaques in AD. In mice, extracellular levels of Aβ in the brain show a clear daily rhythm; levels increase during waking hours and decrease during sleep. Disrupting circadian rhythms in mice by deletion of a key circadian clock gene leads to fragmented sleep timing and loss of Aβ rhythms, suggesting that the circadian system controls Aβ rhythms in the brain, likely by controlling sleep timing. Both sleep deprivation and disruption of circadian rhythms caused accelerated amyloid plaque formation in mice, suggesting a key role for sleep in amyloid plaque regulation. In contrast, increasing sleep by treating mice with an orexin antagonist leads to marked decreases in plaque pathology.

The mechanisms linking sleep and amyloid pathology are not yet known. Because neuronal activity increases Aβ levels, a possible explanation is that rhythms in Aβ levels reflect changes in neural activity during sleep and wake. Supporting this theory is that Aβ rhythms in the brain correlate closely not only with sleep rhythms, but also with lactate levels, which are a marker of neuronal activity. Clearance of Aβ from the brain may also play a role. In mice, sleep is associated with increased lymphatic-like circulation of extracellular fluid through the brain, termed *glymphatic flow* because of the importance of glia in the process. Increased glymphatic flow has been proposed as a mechanism of Aβ clearance during sleep, and ongoing studies in humans are examining the potential role of impaired glymphatic flow in AD (Figure).

The relationship between sleep and circadian rhythms and Aβ seems to be present in humans. Similar to mice, there are daily rhythms in CSF Aβ levels in humans, although it is unknown if the circadian clock regulates these oscillations. Healthy volunteers who experienced acute sleep deprivation for 1 night sustained increases of CSF Aβ levels. Specific disruption of slow-wave sleep for 1 night, with sparing of other types of sleep, also led to increased CSF Aβ levels in healthy volunteers the next morning. Partial sleep deprivation without disruption of slow-wave sleep had no effect on CSF Aβ, emphasizing a key role for slow-wave sleep in regulating CSF Aβ levels. Thus, acute sleep deprivation, and particularly loss of slow-wave sleep, appears to increase Aβ levels in humans, suggesting that chronic sleep deprivation might promote amyloid plaque formation over time. An imaging study using amyloid positron emission tomography suggests that 1 night of sleep deprivation increased amyloid signal in the hippocampus of healthy volunteers. Although this result is controversial and needs replication, it suggests a very rapid effect of sleep loss not only on soluble Aβ levels, but also Aβ aggregation and plaque formation. In general, accumulating evidence from human subjects supports an acute effect of sleep deprivation on Aβ dynamics, while the effect of chronic sleep disturbances on plaque formation is still under investigation.

Therapeutic Considerations

The treatment of sleep and circadian disruption in patients with ADD has been an area of considerable dif-
Sleep and circadian rhythm disruption are emerging as important potential contributors to risk and pathogenesis of AD. Studies in mice and humans support a relationship between sleep and Aβ levels, and mouse studies strongly suggest that sleep and circadian disruption promote amyloid plaque formation (Figure). However, the underlying mechanisms are unclear, and the efficacy of targeting sleep or circadian function for prevention is unknown. The treatment of sleep and circadian symptoms in patients with AD dementia remains challenging, as these patients tend to respond poorly to many typical sleep medicines. Although sleep and circadian rhythms are promising targets for prevention of AD and mitigation of morbidity in patients with symptomatic AD, our understanding of these processes is in its infancy.

Summary

Sleep and circadian rhythm disruption are emerging as important potential contributors to risk and pathogenesis of AD. Studies in mice and humans support a relationship between sleep and Aβ levels, and mouse studies strongly suggest that sleep and circadian disruption promote amyloid plaque formation (Figure). However, the underlying mechanisms are unclear, and the efficacy of targeting sleep or circadian function for prevention is unknown. The treatment of sleep and circadian symptoms in patients with AD dementia remains challenging, as these patients tend to respond poorly to many typical sleep medicines. Although sleep and circadian rhythms are promising targets for prevention of AD and mitigation of morbidity in patients with symptomatic AD, our understanding of these processes is in its infancy.