THEORETICAL REVIEW

Candidate mechanisms underlying the association between sleep-wake disruptions and Alzheimer’s disease

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S U M M A R Y

During wakefulness, extracellular levels of metabolites in the brain increase. These include amyloid beta (Aβ), which contributes to the pathogenesis of Alzheimer’s disease (AD). Counterbalancing their accumulation in the brain, sleep facilitates the removal of these metabolites from the extracellular space by convective flow of the interstitial fluid from the para-arterial to the para-venous space. However, when the sleep-wake cycle is disrupted (characterized by increased brain levels of the wake-promoting neuropeptide orexin and increased neural activity), the central nervous system (CNS) clearance of extra-cellular metabolites is diminished. Disruptions to the sleep-wake cycle have furthermore been linked to increased neuronal oxidative stress and impaired blood–brain barrier function — conditions that have also been proposed to play a role in the development and progression of AD. Notably, recent human and transgenic animal studies have demonstrated that AD-related pathophysiological processes that occur long before the clinical onset of AD, such as Aβ deposition in the brain, disrupt sleep and circadian rhythms. Collectively, as proposed in this review, these findings suggest the existence of a mechanistic interplay between AD pathogenesis and disrupted sleep-wake cycles, which is able to accelerate the development and progression of this disease.

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Regulation of amyloid and tau levels in the brain across the sleep-wake cycle

The aggregation of amyloid beta (Aβ) peptides (predominantly Aβ peptides 1–40 and 1–42; Aβ40 and Aβ42, respectively) into plaques in the brain is a marker of Alzheimer’s disease (AD) and a key component of the ‘Amyloid cascade hypothesis’ [1]. In recent years, increasing evidence has accumulated to support the hypothesis that the production of Aβ peptides in the brain is closely connected to the 24-hr sleep-wake cycle, with high extracellular levels during wakefulness and low extracellular levels during sleep [2–4] (Fig. 1). A major driver for the production of Aβ appears to be neuronal activity, which is higher during wakefulness as compared with sleep. This hypothesis is supported by the observation that unilateral vibrissal stimulation increases, while unilateral vibrissal removal decreases, interstitial fluid (ISF) levels of Aβ in the contralateral barrel cortex of transgenic mice (Tg2576) [5]. In humans, ISF Aβ concentrations have been shown to increase in patients with acute brain damage as neurological status improves, and conversely to fall when neurological status declines [6].

During sleep, the brain remains metabolically and electrically active with preservation of cortico-cortical connectivity during light sleep, i.e., non-rapid eye movement (NREM) sleep stage 1 (N1) and NREM sleep stage 2 (N2) [7–9]. However, a reduction occurs in fronto-parietal functional connectivity with increasing depth of NREM sleep to the point of being significantly reduced in deep sleep [7–11], also called NREM sleep stage 3 (N3) or slow-wave sleep (SWS). Therefore, Aβ production could be postulated to decrease during SWS by virtue of the decreased neuronal activity in this sleep stage. Supporting this hypothesis, cerebrospinal fluid (CSF) Aβ42 levels have been shown to be lowest in humans at around 10:00 h (around 25% lower than peak values), corresponding to a nadir in ISF levels at 04:00 h (as there is a 6-h lag for brain soluble Aβ to reach the lumbar space [12,13]). This represents a time point after which most SWS has typically occurred and after

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which sleep is predominated by sleep stages N1-2 and rapid eye movement (REM) sleep.

Neuropeptides involved in the regulation of the sleep–wake cycle may additionally contribute to the characteristic 24-hr pattern of Aβ peptides in the brain. One such candidate is the hypothalamic neuropeptide orexin-A (hypocretin 1), the level of which increases during wakefulness [14]. A study in transgenic APPswe (Tg2576) mice—a mouse model of AD pathology, which carries the Swedish mutation (K595N/M596L) of the amyloid precursor protein (APP) resulting in higher Aβ peptide levels, and which does not develop behavioral signs of AD—showed that intracerebroventricular administration of orexin at the beginning of the light (i.e., inactive) period could acutely increase both wakefulness and Aβ levels in ISF. Conversely, intracerebroventricular treatment over 24 h with a dual orexin receptor antagonist (almorexant) decreased Aβ ISF levels [2].

Further supporting the role of orexin for Aβ accumulation, daily treatment with almorexant for 8 w reduced the formation of Aβ plaques in several brain regions in APPswe/PS1dE9 mice [2]. In a recent study performed in amyloid transgenic mice in which the orexin gene was knocked out (APP/PS1dE9/OR−/−) [15], loss of orexin resulted in decreased wakefulness and a subsequent reduction in amyloid pathology. In contrast to findings of animal studies, evidence from human studies about the role of orexin in the regulation of Aβ production in the brain is less consistent [16–23]. For instance, a recent study involving patients with the sleep disorder narcolepsy—a disease hallmarked by a progressive loss of brain orexin function [24]—revealed that CSF concentration of Aβ was significantly higher in the patient group with normal CSF orexin-A concentration than in those with low orexin-A concentrations [16]. Moreover, in a separate study, CSF levels of Aβ42 were found to show no relationship in both AD patients and healthy controls [20].

Another key component of AD pathogenesis is the accumulation of intracellular neurofibrillary tangles (NFTs) composed of...
hyperphosphorylated tau (P-tau) protein. Importantly, to date it is not known if the parenchymal levels of tau in the CNS exhibit the same 24-hr rhythmicity as Aβ. Notwithstanding, neuronal activity rapidly increases extracellular tau in mice [27], suggesting that mammalian ISF levels of tau may exhibit a similar neuronal activity-driven 24-hr sleep-wake pattern as Aβ.

Disruptions to sleep and circadian rhythms and the risk of AD

Given the evidence above, an obvious question is: do chronic disruptions to the sleep-wake cycle increase the risk of AD in humans? An increasing number of studies support such a notion, as insomnia [28], self-reported sleep disturbances [29], a decline in sleep duration [30], impaired sleep consolidation [31], delayed or decreased circadian rhythms [32], and sleep disordered breathing (SDB) [33], all have been shown to increase the risk of AD (for a detailed review, see [34]). While AD itself may cause sleep disruptions, the fact that sleep disruptions increase AD risk in non-demented older humans, supports the hypothesis that a chronically disrupted sleep-wake cycle can drive AD pathogenesis. Present evidence from human and animal experiments lends further support to this hypothesis. Experimentally induced sleep disruptions, to date almost exclusively carried out in rodent models, lead to an accumulation of AD-promoting metabolites Aβ and tau in the brain [23,33–37], increase central nervous system (CNS) oxidative stress [38–40], and reduce the structural and functional integrity of the blood–brain barrier (BBB) [41,42], all of which have been hypothesized to promote the development and progression of AD [1,43–44]. Finally, disrupted circadian rhythms, which have been found to even occur prior to the clinical onset of AD [45], have also been linked to neurodegeneration in rodent models [46,47].

Recent results from human and transgenic animal studies also support the idea that AD pathology itself can lead to sleep and circadian disruptions. It has for instance been demonstrated that pathophysiological processes associated with AD, such as Aβ deposition in the brain, alter sleep, as well as disrupt circadian rhythms [3,48,49]. Since Aβ deposition in the brain and cognitive dysfunctions are detectable years prior to the clinical onset of AD [50,51] this suggests that disruptions to the sleep-wake cycle may be a consequence rather than cause of AD pathogenesis. Alternatively, existing evidence leads us to propose that there exists a mechanistic interplay between AD pathogenesis and disruptions to sleep and interrelated circadian rhythms (as illustrated in Fig. 2).

With this in mind, the objective of our review is to systematically frame recent experimental findings from human and animal experiments into a comprehensive overview on candidate mechanisms through which chronically disrupted sleep-wake cycles (e.g., fragmented sleep and circadian disruption) drive AD pathogenesis, and vice versa.

While there are several recent reviews on sleep-wake cycle disruptions and CNS deposition of Aβ peptides [52–56], to our best knowledge no one has yet comprehensively reviewed the recent literature including the role of tau and detailing the wider range of candidate mechanisms that may underlie the harmful association between sleep-wake disruptions and risk of AD.

Candidate mechanisms underlying the association between sleep-wake disruptions and Alzheimer’s disease

Clearance of AD-promoting metabolites from the brain

Soluble Aβ levels are higher in the brain during wakefulness and lower during sleep, indicating that sleep may curb processes leading to Aβ production, concomitantly promoting processes involved in Aβ clearance [2–4]. In contrast, under conditions of acute sleep deprivation, brain Aβ concentrations further increase during the night/inactive period, both in mice and in humans [2,35,57]. In one study, mice were subjected both to acute and chronic sleep deprivation. Acute sleep deprivation during 6 h of daytime (7:00–13:00 h) resulted in around 17% higher ISF Aβ levels in tg2576 mice. Chronic sleep deprivation of APPswe/PS1dE9 mice, which instead lasted for 21 d, more than doubled Aβ levels throughout the brain compared with control animals [2]. Whereas neither Aβ nor tau levels have been assessed in CSF in more long-term sleep deprivation experiments in humans, several studies have confirmed that sleep in humans is associated with lower Aβ levels in CSF. In a study of 26 healthy men (age 40–60y), the 13 individuals who were allowed to sleep for one night showed a 6% decrease in CSF Aβ42 levels across the night; an additional correlation analysis showed that total sleep duration correlated with decreased Aβ42 in CSF — neither of these effects were seen for Aβ40, P-tau or total tau (T-tau) [35]. Meanwhile, the 13 subjects who instead underwent sleep deprivation for one night had no overnight decrease in Aβ42 levels in CSF [35]. With these findings in mind, the question is what mechanism — decreased production or increased clearance — mainly accounts for the drop in brain Aβ levels during sleep, and which of the involved mechanisms is impaired to the greatest extent during sleep loss?

Given the evidence presented above, it has been suggested that sleep may be a major driver for the overnight decrease in Aβ levels in the brain. The clearance of Aβ is driven by local degradation by a wide range of proteases [58], phagocytosis by glial cells [59], egress across the BBB, and Aβ reabsorption through the CSF [60]. Recently, the existence of a CNS paravascular circulation [61] was confirmed and extended by the use of in vivo two-photon imaging in mice [62]. It was determined that the CSF acts in the brain parenchyma like lymph, by flushing out interstitial substances in a process facilitated by glial cells. It was therefore named the glymphatic (a glia-dependent lymphatic) system [62]. Subsequently, another study found that this CSF-ISF exchange increased during sleep in wild-type mice [37], thereby enabling removal of metabolites that typically accumulate during wakefulness, encompassing AD-promoting soluble Aβ peptides. During wakefulness, however, the removal of such metabolites or inert tracers from the brain was not as efficient [37]. The study demonstrated that there was a 60% increase in the interstitial space of brain parenchyma in mice during sleep, as compared with the space found during wakefulness, and sleep was found to increase the convective flow of ISF from the para-arterial to the para-venous space [37] (see Fig. 3). The removal of substances was also greatly increased during sleep, possibly as a result of the concomitant expansion of the interstitial space, evidenced by a doubling in the ratio of Aβ removal from brain parenchyma seen during wakefulness [37]. These effects were furthermore mimicked by infusion of noradrenergic receptor antagonists, suggesting that low adrenergic input is required for this convective clearance to occur. In contrast, awakening of sleeping mice sharply reduced the para-arterial and parenchymal influx of ISF (reduction of ~95%) [37]. This could suggest that sleep disruptions hallmarked by recurrent awakenings, increased time awake after sleep onset, or sleep fragmentation due to SDB, may diminish the ability of the glymphatic system to remove Aβ from the brain. Importantly, a caveat that must be considered is that the study by Xie et al. utilized exogenously administered Aβ to ascertain the clearance function of the glymphatic system during sleep [37]. Thus, it has not yet been shown to which extent the function of the glymphatic system during sleep is of relevance for clearance of endogenously produced Aβ. This also applies to a recent human study, which found MRI-based evidence for modulation of the extracellular space by sleep and wakefulness. Following 24 h of wakefulness combined with cognitively demanding task, increased subcortical but not global gray and white matter volumes were
observed in the healthy participants, as well as decreased volume of the brain ventricles, compared with the volumes observed after normal sleep. These changes reverted after recovery sleep. However, any relationship to a possible glymphatic system or potential increased clearance of metabolites during human sleep, as compared with wakefulness, was not examined [63].

While aggregation of P-tau into NFTs represents the other main pathologic feature of AD [64], clearance of tau has not been well characterized. The protein exists in six isoforms, ranging from 352 to 441 amino acids (ten times the size of Aβ peptides), without any known active transport mechanisms to blood like Aβ. CSF bulk flow and in situ degradation are the most likely mechanisms for clearance of tau released into the interstitial space, but a recent study in mice showed that extracellular tau could also be cleared from the brain via the glymphatic pathway under anesthesia [65]. To the best of our knowledge, no study has directly investigated the role of sleep in the degradation or removal of tau protein from the ISF, and whether its removal can be impaired by sleep loss. One clinical study has shown that CSF P-tau or T-tau levels are not affected by one night of sleep loss in healthy individuals [35], although as noted by the authors, the long turnover of tau (11 d; [27]) would likely have required a considerably longer sleep loss paradigm to study how these dynamics are affected by disrupted sleep.

Because sleep clears cellular waste from the extracellular space of the brain parenchyma [37], it is nonetheless likely that sleep promotes tau protein removal from the brain ISF and reduces its aggregation into NFTs. Suggestive of this, studies using mice that develop AD pathology (3Xtg; produce amyloid plaques and NFTs) suggest that altered sleep-wake patterns can increase tau levels in the brain [36,66]. In one of these studies, six weeks of sleep restriction for 6 h/day increased cortical Aβ and P-tau to about 2-fold of control animals, although the changes were not significant [66]. In another study, two months of sleep disruption by prolonged daily light exposure (20/4-h light—dark cycle) led to a greater than 50% significant increase of the insoluble fraction of tau in the brain, as compared with control mice that were maintained on a 12/12-h light—dark cycle [36].

As reviewed in this section, the recent finding that sleep promotes the function of a glymphatic system, resulting in enhanced removal of AD-promoting metabolites, is intriguing, but still requires additional research. For instance, it is still unclear whether sleep-driven clearance of metabolites is also present in humans and whether it can increase the clearance rate of neurotoxic substances to the same extent as in mice. Moreover, further research is needed to disentangle the contribution of normal and disrupted sleep stages and circadian mechanisms to this function, for example to address whether interventions that enhance specific sleep stages can improve the clearance of metabolites from the brain in subjects genetically prone to develop AD. Given that activity profiles of neurotransmitter and neuropeptide systems vary across sleep stages (e.g., high cholinergic activity during REM sleep vs. low cholinergic activity during SWS [67]) further studies are needed to ascertain how sleep stage-specific neurotransmitter and neuropeptide patterns contribute to the clearance of AD-promoting metabolites from the ISF. A recent study demonstrated that when old (18-month-old) and young (2- to 3-month-old) mice were compared, the older mice exhibited a marked reduction in the CSF-ISF exchange, with a 27% reduction in the pulsatility in arterioles in the brain, and a loss of perivascular aquaporin-4 (AQP4) polarization [68]. Furthermore, the older mice exhibited a 40% reduction of their ability to clear Aβ that had been injected intraparenchymally [68]. However, no age-associated decline was seen in the ability of sleep to increase the interstitial space. This suggests that deterioration of this part of the glymphatic system does not contribute to the age-associated increase in the risk of AD, to which disrupted sleep may contribute. However, the influence of sleep deprivation

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**Fig. 2.** Overview of proposed mechanisms through which disruptions to the sleep-wake cycle form a positive feedback loop with AD pathogenesis in humans. Abbreviations: Aβ, amyloid beta; AD, Alzheimer disease; CNS, central nervous system; BBB, blood–brain barrier; EE, energy expenditure; NFTs, neurofibrillary tangles.
on the glymphatic system in young compared with old mice was not investigated and as such warrants further investigation.

**CNS oxidative stress**

In addition to the accumulation and deposition of Aβ and NFTs in the brain, CNS oxidative stress resulting from increases in reactive oxygen species (ROS) and reactive nitrogen species (RNS) has also been proposed to promote the development and progression of AD [43]. In APPswe mice, oxidative stress precedes deposition of Aβ [69], and evidence of oxidative damage has been observed in human AD subjects studied postmortem, with the greatest pathology found early in the disease [70]. Defective mitochondria, which are a major source of ROS, are seen in AD as well as in what is the greatest risk factor for AD, i.e., aging, and may contribute to AD pathology by altering how APP and tau proteins are processed [71]. Conversely, APP and Aβ have been shown to interfere with mitochondrial functions and enzymes [71], in what may develop into a positive feedback loop. ROS and RNS play essential roles under physiological conditions (e.g., induction of host defense). However, excessive production of reactive species can induce cellular stress through lipid peroxidation and protein oxidation. Once initiated, such oxidative processes can lead to damage to vital cellular components such as proteins, lipids, and nucleic acids, which can finally result in cellular death [43].

Several animal studies have found loss of sleep to increase oxidative stress in the brain [38,39,72,73], suggesting it might be a candidate mechanism underlying the association between sleep loss and AD. For instance, in one such study in mice [72], three nights of extended wakefulness (via environmental enrichment during the rest cycle) increased signs of oxidative stress in locus coeruleus (LC) neurons, as reflected by increased production of superoxide. Another study found that sleep loss in mice (for 72 h using a multiple platform method) increased oxidative stress in the hippocampus and was linked to learning deficits, as both the oxidative stress and the learning deficits could be prevented if anti-oxidative agents (N-tert-butyl-alpha-phenylnitrone, vitamin E or melatonin) were administered prior to the sleep deprivation [73], supporting a detrimental role for oxidative stress following even short term sleep loss. It must however be noted that other studies in mice have not found signs of increased oxidative stress in the cortex using total sleep deprivation paradigms [74] or following REM sleep deprivation after which whole-brain extracts have been analyzed [75]. One explanation for these discrepant results could be that oxidative stress is primarily apparent in brain regions that fire at increased rates across sustained wakefulness, comprising the LC [72]. Supporting this notion, such brain regions appear to be more sensitive to cellular damage that can lead to cellular death following sleep disruptions in humans with and without AD [76,77].

An important question that remains unanswered is how can sleep loss cause oxidative stress in the brain? One explanation could be that cellular scavenger mechanisms against reactive species are compromised in function. Supporting this notion, three nights of extended wakefulness were shown to reduce the activity of sirtuin type 3 (SirT3) in the mouse brain [72]. SirT3 is a nicotinamide adenine dinucleotide-dependent enzyme that is localized to the mitochondrial membrane and which upregulates many antioxidant defenses. Anti-oxidative mechanisms may also decrease in certain brain regions following sleep loss, especially in older animals [78], which also display lower CNS levels of defensive mechanisms such as lower levels of SirT3 [79].

**Blood-brain barrier (BBB) integrity**

Neurons demand a nearly continuous supply of energy metabolites, as they have only limited energy reserves. This requires a continuous metabolite exchange between circulating plasma and the brain. This high rate of molecular exchange, however, also implies that neurons are exposed to many potentially harmful factors derived from the periphery. To prevent the brain from uncontrolled entry of blood factors and toxins as well as an unfavorable efflux of CNS metabolites into the periphery, the BBB consists of endothelial cells lining brain capillaries that tightly regulate the flow of nutrients, ions, and fluids between both compartments. As reviewed in Ref. [44], BBB dysfunctions (e.g., endothelial loss and loss of tight junction proteins) have been proposed to contribute to the...
development and progression of AD, as they impair Aβ clearance from the brain, lead to increased influx of circulating Aβ into the brain, and elevate expression and processing of the Aβ precursor protein.

Importantly, emerging evidence suggests that sleep disruption may impair the function of the BBB [41,42]. For instance, six days of sleep restriction in mice, resulting in a mild 13% increase in total wake time, led to a reduced expression of tight junction proteins by BBB endothelial cells [42]. This reduction was paralleled by increasing paracellular permeability of the BBB to small substances, which under physiological conditions mainly reach the brain via a saturable transport system located at the BBB [42]. The expression of glucose transporter 1 (GLUT1), a protein that mediates glucose uptake through cerebral BBB micro-vessels, was also reduced following the sleep restriction paradigm. Importantly, functional and structural alterations to the BBB have also been found in AD patients, including reduced expression of GLUT1 [80–82]. Further highlighting the importance of glucose transport, a recent study found that GLUT1-deficient (Slc2a1<sup>−/−</sup>) mice with APPswe expression had increased BBB permeability, reduced dendritic spines and cognitive deficits [83].

As the increase in paracellular permeability following six days of sleep restriction in mice returned to baseline after 24 h of recovery sleep [42], this however suggests that impaired BBB integrity following short periods of sleep restriction is a reversible process. It is currently not known whether sleep restriction-induced BBB disruptions may accelerate neurodegenerative processes involved in AD, such as Aβ plaque formation. This appears especially relevant given that a recent study found that preclinical AD mouse models (e.g., PS2-APP and hTauP301S) do not per se display disrupted BBB function [84], as assessed by passive antibody uptake into the brain. This could suggest that the role of the BBB may be dissociated from the pathogenic burden posed by Aβ accumulation on e.g., circadian rhythms and sleep patterns, whereby BBB disruption (due to e.g., sleep restriction) may increase AD pathology, but not vice versa.

Circadian disruption

Accumulating evidence connects disruptions of circadian rhythms, as frequently found in AD patients [45], but also in shift workers who are also more likely to be short sleepers [85], to neurodegeneration and cognitive aging [86,87]. For instance, one study found that shift workers without the ability to recover for longer time periods between demanding periods of shift work (less than five versus over 14 d of recovery time) had increased temporal lobe atrophy observed on magnetic resonance imaging [86]. In addition, a 5-year prospective study of 1282 women found that delayed peak of activity rhythms, and decreased amplitude and robustness thereof, conferred an increased risk of mild cognitive impairment (MCI) or dementia [32].

Recent studies have begun delineating how disrupted circadian rhythms may contribute to neurodegeneration [summarized in [46] and [47]]. Mice with ablated Clock or Bmal1 – genes regulating central and peripheral molecular clocks – exhibit impaired sleep parameters. Homozygous Clock mutants sleep approximately 2 h less than wild-type mice [88], and they exhibit increased signs of oxidative stress in the brain [89,90]. Neuronal- and glial-specific deletion of the master clock gene Bmal1 in mice also increased neurodegeneration, as evidenced by degeneration of synaptic terminals and impaired cortical functional connectivity, as well as neuronal oxidative damage and impaired induction of several redox defense genes, such as Adh2 and Nqo1 [90]. This was observed even though such genetically targeted ablation (using a Nestin-Cre driver) does not fully abolish expression of these clock genes in the suprachiasmatic nucleus (SCN) – the pacemaker clock that entrains other circadian oscillators – and accordingly activity and sleep rhythms remained largely unaffected [90]. Similarly, disrupting the genetic clock machinery via knockout of Bmal1 or Per2 has been shown to impair murine hippocampal neurogenesis by perturbing the conversion of quiescent neural progenitor cells into newborn neurons in the mouse hippocampus [91].

Although these studies establish a connection between circadian disruption and neurodegeneration, which may be linked to AD, they leave several gaps in our knowledge warranting further investigation. For instance: does glial–specific deletion of master clock genes (e.g., Bmal1) alter the ability of the glymphatic system to remove AD-promoting metabolites from the brain parenchyma, and as an extension, to what extent does the glymphatic system depend on properly aligned circadian rhythms? Circadian rhythm disturbances are frequently seen even in patients with preclinical AD [45]. The possible contribution of circadian disruption to AD disease progression has been showcased by a study in which bright- vs. low-light exposure in combination with melatonin vs. placebo was used to study the effect of synchronizing circadian rhythms in over 189 patients, 63% of whom had probable AD, over an average follow-up period of 15 months. Both bright light and melatonin had positive effects: the light therapy was for example able to slow cognitive decline (as measured by the mini-mental state examination) [52]. Thus, it may also be worth investigating if therapeutic synchronization of the circadian timing system (e.g., through bright light therapy) can help re-establish circadian Aβ dynamics in ISF and CSF in humans suffering from circadian rhythm disorders or at increased risk of AD (e.g., ApoE4 carriers), and if such therapies can curb the development and progression of this disease.

Notably, functional weakening of the circadian system, characterized by phase advance, increased fragmentation and reduced amplitudes of circadian rhythms, is a well-documented consequence of aging [46]. Given that synchronized activity of multiple circadian clocks in the brain has been suggested to be critically important for a number of CNS processes [46], it could be speculated that loss of phase coherence between these CNS clocks due to normal aging, promotes neurodegenerative processes associated with AD pathology. Supporting this view, it has been shown that, when comparing young and elderly adults who were either positron emission tomography (PET) amyloid positive or negative, 24-h fluctuations in CSF Aβ levels decreased with age and even more in those who were ‘amyloid positive’ [2,57]. The loss of the dynamic pattern was more pronounced for Aβ42 than for Aβ40, most likely due to its greater propensity to aggregate in amyloid plaques [93].

Disruptions to the sleep-wake cycle as a consequence of AD-related processes

Multiple lines of evidence suggest that neurodegenerative processes associated with AD can cause sleep and circadian disruptions in humans. This is particularly relevant to the interpretation of findings from epidemiological studies investigating the association between sleep disruption and AD features, as they do not typically account for the occurrence of preclinical AD in a substantial proportion of cognitively healthy older individuals who are included in such studies [50,51]. For instance, a recent study involving 45 older adults (12 with AD) demonstrated that individuals with AD had fewer intermediate nucleus neurons than controls at the time of death [76]. The intermediate nucleus is considered as the human homolog of the rodent ventrolateral preoptic nucleus, a brain region that promotes sleep by inhibiting wake-promoting brain regions, which include the lateral hypothalamic area, raphe nucleus, tuberomammillary nucleus and the LC (for more details concerning the neurobiology of sleep regulation, please see [94]). Additional support for the existence of a bidirectional rather than unidirectional link between sleep disruption...
and AD has been provided by a recent study involving 26 cognitively healthy (i.e., non-demented) elders. In this study, it was revealed that those with high Aβ burden in the medial prefrontal cortex (a typical feature of AD) had lower slow-wave activity (SWA) during NREM sleep. Furthermore, prefrontal Aβ burden was found to be associated with impaired sleep-dependent memory consolidation, most likely mediated through its effects on NREM SWA [48]. Adding further support to the hypothesis that amyloid deposition may alter sleep characteristics in the preclinical stage of AD, three additional studies performed in younger-old and middle-old cognitively healthy individuals have reported shorter sleep duration and/or lower sleep quality to be associated with greater CNS amyloid burden [94–97].

Collectively, evidence from both human and animal studies provide a strong rationale for hypothesizing that poor sleep and disrupted circadian rhythms may be a potential early marker of neuropathology during the long preclinical phase of AD.

Conclusions

Observational studies have found that patients with insomnia or sleep disruptions in mid-life to old age have an increased risk of pathological changes that precede AD (e.g., CNS increased amyloid burden, neurofibrillary tangles), as well as an increased risk of dementia and AD. In line with these findings, experimental studies have demonstrated that sleep disruptions result in higher levels of markers that are associated with AD, increase CNS oxidative stress, can damage the blood–brain barrier in mice, and disrupt clearance of AD-promoting Aβ peptides. Importantly, patients with impaired cognition or increased Aβ burden also show signs of impaired sleep, and animal models have shown that Aβ deposition can directly drive impaired sleep. Collectively, current evidence points toward the existence of a mechanistic loop between AD pathogenesis and disrupted sleep and circadian rhythms.

Given the alarming increase in the number of people who are afflicted by chronic sleep problems [108] and AD pathology [109], studies with long follow-up periods and repeated observations initiated prior to the clinical onset of AD are needed to further disentangle the contribution of sleep and AD pathology in their intertwined relationship. The ideal longitudinal studies will assess sleep and circadian rhythms both subjectively and objectively (e.g., by actigraphy and EEG-based sleep monitoring), simultaneously with AD biomarkers (CSF biomarker levels, imaging using PET and, potentially, of glymphatic flow) and AD- and sleep-modulating risk factors, such as genetics, co-morbidities, exercise and exposure to environmental light and stress. The extent to which general improvements in sleep, or e.g., targeted sleep-stage enhancement, can help in lowering the risk or reversing signs of accelerated cognitive ageing, MCI/AD, or other neurodegenerative diseases represent an important parallel path of research in this emerging interdisciplinary field.

Practice points

Disrupted sleep has been found to be associated with an increased risk of Alzheimer’s disease (AD) through several mechanisms:

1. Observational studies have found that patients with e.g., insomnia or sleep disruptions in mid-life to old age have an increased risk of pathological changes that precede AD (e.g., CNS increased amyloid burden, neurofibrillary tangles), as well as an increased risk of dementia and AD.

2. Sleep restriction results in higher levels of markers that are associated with AD, increases CNS oxidative stress and can damage the blood–brain barrier in mice.

3. Clearance of the AD-promoting Aβ peptides is greatly enhanced during sleep in mice and awakening the mice disrupts this process.

4. Importantly, patients with impaired cognition or increased Aβ burden also show signs of impaired sleep, and animal models have shown that Aβ deposition can directly drive impaired sleep, in what can turn into a positive feedback loop.
Research agenda

Future research should address the following questions with regards to sleep and AD pathogenesis:

1. The contribution of sleep disruptions to AD pathogenesis, and of AD pathogenesis to sleep disruption in humans, by conducting long follow-up studies that are initiated prior to the presence of significant amyloid and tau burden in the brain that may otherwise conceal the contribution of each factor.
2. How clearance of metabolites in the CNS is regulated in humans and what sleep stage(s), circadian and underlying neurobiological mechanisms are the greatest contributors to this function.

The extent to which general improvements in sleep, or e.g., targeted sleep-stage enhancement, can help in lowering the risk or reversing signs of accelerated cognitive aging, MCI/AD, or other neurodegenerative diseases.

4. The impact of genetic variants in influencing the extent by which sleep disruption can confer an increased risk of AD.

Conflict of interest

The authors do not have any conflicts of interest to disclose.

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