

Burden of Insomnia and Sleep Disturbances and the Impact of Sleep Treatments in Patients with Probable or Possible Alzheimer's Disease: A Structured Literature Review

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Accepted 3 December 2021

Pre-press 1 February 2022

Abstract.

Background: Sleep disturbances are frequent in Alzheimer's disease (AD). This review summarizes the impact of sleep disturbances on AD patients and their caregivers and the effects of currently available sleep therapies.

Methods: Published studies (January 1985–March 2020) assessing the burden associated with insomnia/sleep disturbances in the AD population and insomnia treatment effects were identified by searching PubMed, Embase and Cochrane Library and screened against inclusion criteria.

Findings: This review included 58 studies which assessed patient and caregiver burden, institutionalization, and insomnia treatments in AD patients with sleep disturbances. Sleep disturbances were associated with worse cognition, functional ability, and behavioral and neuropsychological functioning, including increased depression and anxiety. Health status and quality of life of both patients and caregivers were reduced in the presence of sleep disturbances, with caregiver burden driven largely by disruptive nocturnal behaviors including nighttime awakenings and wanderings. Sleep disturbances were also associated with institutionalization. Although significant associations between sleep problems and clinical outcomes were apparent, there was generally no control for other influencing factors (e.g. cognitive status). With respect to insomnia treatments, bright light, and behavioral therapies as well as drugs such as trazodone, risperidone and suvorexant showed some promise in AD patients, but studies were primarily small and limited data were available, particularly in regard to insomnia treatment effects on associated clinical burden.

Interpretation: Sleep disturbances are a significant problem for AD patients and caregivers. They are associated with behavioral and psychological problems and cognitive decline and impose a burden on caregivers, but remain poorly characterized and under-researched. As the global population is aging and AD is on the rise, data from larger, prospective trials are required to fully understand the clinical correlates of sleep disturbances and the impact of insomnia treatments on AD patients and their caregivers.

Keywords: Insomnia, Alzheimer's disease, sleep disturbances, literature review, caregiver burden, clinical burden, institutionalization, treatment guidelines

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INTRODUCTION

Alzheimer's disease (AD) is a progressive neurodegenerative disease, characterized by continuing cognitive decline and memory loss; [1] it accounts for 60% to 80% of dementia cases in adults > 65 years of age. With an aging global population, the AD population is expected to triple over the next 40 years, to over 13 million cases [2].

AD is associated with significant healthcare costs and resource utilization, particularly when patients require admission to care facilities [3]. The behavioral symptoms of dementia, including disturbed sleep, nighttime awakenings and wanderings place a substantial burden on family and caregivers, and lead to the early institutionalization of AD patients [2, 4].

Insomnia is characterized by difficulty initiating or maintaining sleep among individuals with adequate opportunity to sleep and is associated with daytime consequences, such as an inability to perform daily activities [5]. Insomnia symptoms and sleep disturbances (collectively referred to in this manuscript as sleep problems) increase with age and are common in AD patients, with a prevalence of >50% [6–8]. AD patients also experience more severe symptoms of insomnia and sleep disturbances, such as increased sleep latency and reduced sleep maintenance, as well as progressive deterioration and instability of circadian rhythms, compared to older adults without AD [9, 10]. Moreover, insomnia is also a risk factor for AD, [11] and a bidirectional relationship is thought to exist between poor sleep and AD pathology [12].

Despite the high prevalence and substantial disease burden of insomnia in AD patients, including negative impacts of insomnia symptoms on mental and physical health, quality of life (QoL) and functional ability, treatment options for this patient population are limited. Benzodiazepine receptor agonists are associated with an increased risk of falls, fractures, and clinically significant impairments in balance and cognition upon awakening [13]. The American Geriatrics Society strongly recommends against the use of benzodiazepines and related non-benzodiazepines in elderly patients [14]. Moreover, current non-pharmacological treatments such as cognitive behavioral therapy for insomnia (CBT-I) may not be effective as AD patients may be unwilling or unable to participate in therapy [5]. Due to the limitations of non-pharmacological and pharmacological treatments for the management of insomnia in AD patients there is a significant unmet need. Currently,

one medication (suvorexant) has been specifically approved for the treatment of insomnia in AD patients by the US Food and Drug Administration (FDA).

This literature review aims to summarize the clinical, economic, and QoL impact of insomnia and sleep disturbances on patients and their caregivers, as well as the abilities of currently available therapies to treat sleep problems and reduce their impacts in this population.

METHODS

Search strategy

We searched Publisher Medline (PubMed), Excerpta Medica Database (Embase) and Cochrane Library for articles published between January 1985, and March 2020, using combinations of keywords and Medical Subject Headings (MeSH) terms pertaining to AD, insomnia/sleep disturbances, disease and caregiver burden, healthcare resource utilization (HCRU), treatment patterns and guidelines (Table S1 through Table S9 in the supplementary appendix). Reference lists of included publications were reviewed to identify relevant publications not captured through the database searches.

Criteria for study inclusion/exclusion

Eligible studies and reviews met the following inclusion criteria: (1) AD patients with insomnia, or sleep disorders, disturbances or dysfunctions (specific sleep disorder/disturbance not reported); (2) disease burden (QoL, functional status, cognitive functioning, behavioral symptoms, and impact on daily living) and caregiver burden outcomes, HCRU, treatment patterns and guidelines; (3) peer-reviewed; (4) English language. Publications were excluded if they included only sleep disorders other than insomnia such as hypersomnia, parasomnia, excessive daytime sleepiness, and obstructive sleep apnea. Where studies were published as both research articles and conference abstracts, the latter were excluded. Full PICOS criteria are reported in the appendix (Table S10).

Data extraction and synthesis

Data extraction was performed by one reviewer and cross-checked by a second reviewer using standardized forms to ensure accuracy. Data from the extracted publications were qualitatively synthesized to summarize the findings of the literature review.

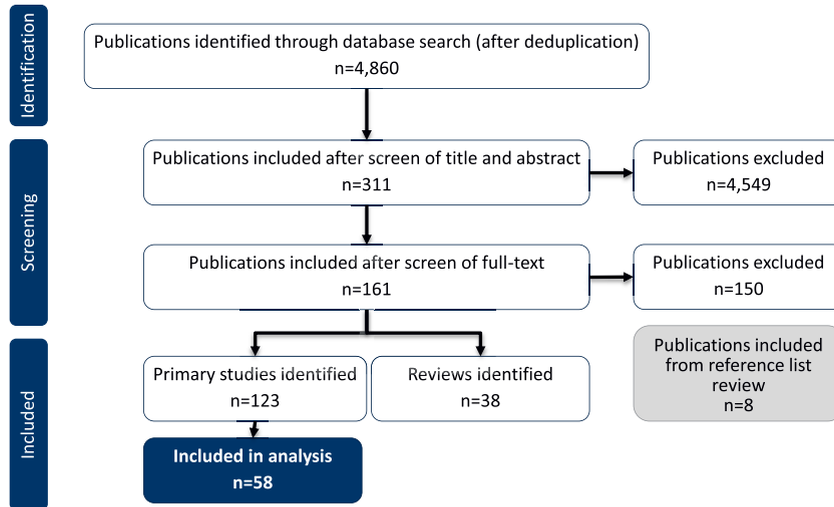


Fig. 1. Study flow diagram.

RESULTS

The search identified 4,860 articles, from which 311 were taken forward to full text review. After screening the full texts against the eligibility criteria, 161 publications were included (123 primary studies and 38 reviews). Eight more publications were included through an additional review of the reference lists (Fig. 1). The majority of publications were excluded for not reporting data for AD patients with insomnia/sleep disturbances.

Only studies characterizing the clinical and caregiver burden of sleep disturbances in patients reported to have AD (including probable or possible AD) were included in this review (58 studies). Clinical burden included the impact of sleep disturbances on patients' health and wellbeing. Studies reporting prevalence and incidence data for sleep disturbances in AD patients and those reporting data for mild cognitive impairment (MCI) were not included in this review (65 studies). To avoid duplication, only primary studies are included in the results. Relevant reviews identified have been included in the Discussion.

The majority of included studies reported data for non-specific sleep disturbances in AD, with fewer evaluating insomnia specifically. Table 1 summarizes the included studies and sleep problem definitions.

Clinical burden of sleep disturbances in AD and impact on QoL

Sleep disturbances were not consistently characterized across studies. Many relied on the

Neuropsychiatric Inventory (NPI), which has a single item to screen for the presence of sleep disturbed behaviors, that incorporates symptoms of awakening the caregiver at night, waking early in the morning and daytime napping, and rates the frequency, severity and distress [4, 29, 72]. An important caveat is that there is insufficient clinical description of the nature of the sleep problems in most studies, and there is not a standardized tool for assessing sleep in AD patients. Despite this limitation, sleep problems in AD patients were associated with impairments across several domains (Table 2). No studies specifically assessed the clinical burden of insomnia in AD.

Cross-sectional studies found significant negative correlations between Mini-Mental State Examination (MMSE) scores and sleep disturbances, suggesting that sleep disturbances are associated with worse cognition [29, 71]. Similar associations were reported for patients' functional impairment [4, 19, 28, 29, 51, 71]. Associations between sleep disturbances and decreased cognitive functioning were reported in five of the nine studies identified (Table 2) [23, 25, 28, 29, 54, 58, 63, 70, 71]. Similarly, associations between sleep disturbances and daily functioning were reported in 10 of the 14 studies identified (Table 2) [4, 19, 28, 29, 47, 51, 58, 68, 70, 71]. Even studies that did not find an association between cognition or functional status and sleep problems (e.g., de Oliveira et al., and Moran et al.) reported sleep problems associated with behavioral disturbances, as defined by the NPI or BEHAVE-AD [25, 54]. Moran et al. found no significant cognitive or functional impairment differences between

Table 1
Included studies and definitions used for cognitive impairment and sleep problems

Study	Country	Setting	Sample size	Diagnosis	Definition of sleep problem
Allegrì, 2006 [15]	Argentina	Clinic/hospital (19 patients were institutionalized)	82	Probable or possible AD	Sleep disturbances, determined using a Spanish version of the NPI (unusual behavior at nighttime)
Ancoli-Israel, 2003 [16]*	US	Nursing home	92	Probable or possible severe AD	Studied effects of treatment on sleep and circadian activity rhythms, measured via wrist actigraphy
Back, 2019 [17]	South Korea	National Health Insurance Service	2,796,871 insomnia patients of whom 138,270 had newly diagnosed AD	Probable or possible AD or VD	Diagnosed insomnia (ICD-10 codes)
Bianchetti, 1995 [18]	Italy	Discharged from dementia unit	86	Probable or possible AD	Insomnia recorded at discharge (not defined)
Bliwise, 1995 [19]	US	Special AD care unit	47	Probable or possible AD	Moderate and severe sleep disturbances defined by ≥ 1 or ≥ 2 observations of wakefulness during night, respectively
Brusco, 1999 [20]*	Argentina	Outpatients	41 (retrospective analysis including only AD: 14)	Probable or possible AD or VaD	Sleep disorder/sleep disturbances reported via sleep logs (caregiver-reported)
Camargos, 2014 [21]*	Brazil	Community/outpatients	30	Probable or possible AD	Sleep disturbances: nighttime insomnia/sleep complaints reported after AD diagnosis, ≥ 2 sleep disorders on NPI.
Camargos, 2015 [22]	Brazil	Hospital outpatients	41	Probable or possible AD	Sleep disturbances causing emotional distress to caregivers (score of ≥ 1 on NPI; caregiver-reported) and sleep profile determined via actigraphy
Chew, 2019 [23]	Singapore	Hospital memory clinic	64	Probable or possible AD with MCI	Sleep duration, defined by the PSQI
Colenda, 1997 [24]*	US	Community-based	5	Probable or possible AD	Disturbed circadian rest-activity cycles (caregiver-reported frequent nighttime awakenings and daytime napping)
de Oliveira, 2014 [25]	Brazil	Outpatient neurology clinic	217	Probable or possible AD	Unsatisfied sleep defined by patient (confirmed by caregiver)
Donaldson, 1998 [26]	UK	Home	100	Probable or possible AD	Sleep disturbances rated using a subsection of the MOUSEPAD (caregiver-reported)
Dowling, 2005 [27]*	US	Care facility	46	Probable or possible severe AD	Rest-activity disruptions, including insomnia, frequent nighttime awakenings, wandering at night, unusually early morning awakenings, "sundowning," and excessive daytime sleepiness
Escudero, 2019 [28]	Colombia	Hospital-based memory clinic	47	Probable or possible AD	Sleep problems defined using the CUSPAD and NPI
Garcia-Alberca, 2013 [29]	Spain	Outpatient clinic	125	Probable or possible AD	Sleep disturbances determined using the sleep disorders item from the NPI; caregiver-reported
Gehrman, 2009 [30]*	US	Nursing homes	41	Probable or possible AD	Various sleep parameters assessed actigraphically
Gehrman, 2018 [31]	US	Clinic/center	130	Probable or possible AD	Sleep disturbances defined by caregiver ratings; defined as a sleep latency > 30 min, ≥ 3 nocturnal awakenings per night, frequency of nocturnal wandering > 1 per month, frequency of loud snoring > 2 per week, and/or spending > 1 h asleep in day

Grace, 2000 [32]	UK	Clinic/hospital	37 (20 with AD)	Probable or possible AD or DLB	Sleep disturbances defined using the ESS and PSQI
Hart, 2003 [33]	NR	Memory clinic	100	Probable or possible AD	Sleep disturbances defined using the NPI (caregiver-reported)
Hannesdottir, 2013 [34]*	US	NR	81	Probable or possible AD and MCI	Not defined; study assessed sleep improvements in AD
Herring, 2020 [35]*	Canada, Finland, Italy, New Zealand, Peru, South Korea, UK, US	Memory/research clinics	277	Probable or possible AD	Insomnia; meeting DSM-5 diagnostic criteria for insomnia and confirmed by mean total sleep time < 6 h over screening and baseline PSG (neither night > 6.5 h)
Ishikawa, 2016 [36]*	Japan	Hospital/clinic	12	Probable or possible AD	Sleep problems/sleep architecture changes (subjective and objective measures)
Kabeshita, 2017 [37]	Japan	Seven psychiatry and neurology centers	684	Probable or possible AD	Sleep disturbances defined using the NPI
Kazui, 2016 [38]	Japan	Outpatients from multiple centers/hospitals	1,598 (1,091 with AD)	Probable or possible AD, DLB, VaD, and FTLD	Sleep disturbances defined using the NPI (caregiver-reported)
Kim, 2017 [39]	NR	NR	46	Probable or possible AD (early stage)	Sleep problems defined using the NPI
La, 2019 [40]*	NR	Memory center	50	Probable or possible AD, MCI or non-impaired	Sleep disturbances (absent or present) documented by a physician; insomnia defined separately
Lebrija, 2016 [41]	Mexico	Hospital/clinic	64	Probable or possible AD	Sleep disturbance defined using the SDI and NPI
Lee, 2018 [42]*	Korea	Home (study specified that 'home-based' light treatment was used)	10	Mild or moderate probable or possible AD	Insomnia symptoms ≥ 3 times per week and/or PSQI score of ≥ 5
Leger, 2017 [43]	France, Germany, Spain, Italy, Portugal, Poland, US, Canada, Australia	Outpatients	208	Mild to moderate probable or possible AD	Sleep disturbances (measured by actigraphy)
Leng, 2020 [44]	China	Outpatient geriatric clinic	221 (AD: 17, MCI: 67, subjective cognitive decline: 91, non-impaired: 46)	Probable or possible AD, MCI, subjective cognitive decline, non-impaired	Sleep disturbances defined using the PSQI. Assessed over four weeks, based on a 0–3 scale for each item
Lukovits, 1992 [45]	US	Long-term care facility	33	End stage probable or possible AD	Sleep disturbances defined via caregiver questionnaire
McCarten, 1995 [46]*	US	Clinic ward	7	Probable or possible AD	Sleep disruption; caregivers' complaint that patients were frequently up at night

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Table 1
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Study	Country	Setting	Sample size	Diagnosis	Definition of sleep problem
McCurry, 1999 [47]	US	Community-based	205	Probable or possible AD	Sleep problems defined by the rating of seven different sleep behaviors from caregivers
McCurry, 2003 [48]*	NR	Community-based	22	Probable or possible AD	Sleep disturbances (≥ 1 sleep problem on the NPI occurring ≥ 3 times a week)
McCurry, 2004 [49]	US	Community-based	153	Probable or possible AD	Sleep disturbances: multiple awakenings at night (caregiver-reported)
McCurry, 2005 [50]*	US	Community-based	36	Probable or possible AD	Sleep problems (≥ 2 sleep problems on NPI nighttime behavior scale occurring ≥ 3 times per week; caregiver-reported)
McCurry, 2006 [51]	US	Community-based	46	Probable or possible AD	Sleep problems; ≥ 2 sleep problems on the NPI that occur $\geq 3-6$ times a week (caregiver-reported), objective sleep measures also used in study (actigraph)
McCurry, 2011 [52]*	US	Community-based	132	Probable or possible AD	Sleep problems; ≥ 2 sleep problems occurring several times a week, measured by the 7-item SDI
Meguro, 2004 [53]*	Japan	Nursing home	34	Probable or possible AD	Disturbed sleep/wake patterns with wandering; patients manifested wandering behavior or aggressiveness for more than 4/7 days (caregiver-reported via written criteria, and BEHAVE-AD)
Moran, 2005 [54]	Ireland	Recruited from memory/aging center	224	Probable or possible AD	Sleep disturbances, defined using the question about diurnal rhythm disturbance on the BEHAVE-AD questionnaire (informant-reported)
Mulin, 2011 [55]	US	Recruited from memory/AD centers	103	Mild or moderate probable or possible AD	Sleep disturbance measured via actigraph and caregiver reports
Okuda, 2019 [56]	Japan	Recruited from Japanese register	496	Mild to moderate probable or possible AD	Clinically diagnosed insomnia: 30.4%; circadian rhythm sleep disorder: 22.2%; parasomnia: 7.9%; narcolepsy: 5.4%; sleep apnea: 5.4%; other sleep difficulties: 5.4%; no diagnosis: 49.4%; and SDI (Japanese version)
Ownby, 2010 [57]	US	Clinic	395	Probable or possible AD	Sleep problems (caregiver-reported)
Ownby, 2014 [58]	US	University-based memory disorder clinic	344	Probable or possible AD	Sleep disturbance (caregiver-reported)
Pang, 2002 [59]	US, Taiwan, Hong Kong	Clinic/hospital/research centers	289 (US: 169; Taiwan: 89; Hong Kong: 31)	Probable or possible AD	Sleep problems defined using the NPI (caregiver-reported)
Petrescu, 2019 [60]*	Romania	Inpatients	43	Probable or possible AD	Sleep disturbances/insomnia symptoms defined from subjective patient and family caretaker interview and objective nursing sleep log reviews
Ribeiro, 2018 [61]	Brazil	Outpatient neurology clinic	NR	Mild and moderate probable or possible AD	Sleep disturbances, defined using AIS and ESS

Scoralick, 2017 [62]*	Brazil	Community	24	Probable or possible AD	Sleep disorders established based on caregivers' emotional distress per the sleep and nighttime behavior item of NPI (score ≥ 2)
Shin, 2014 [63]	South Korea	Dementia clinic	117 (63 AD and 54 matched non-demented controls [including 42 with MCI])	Probable or possible AD	Subjective sleep problems assessed via the Korean version of the PSQI (problems present during the previous month)
Simoncini, 2015 [64]*	Italy	Nursing homes	129	Probable or possible AD and MCI	Primary and secondary insomnia defined using the PSQI
Singer, 2003 [65]*	US	Clinics and long-term care facilities	157	Probable or possible AD	Nighttime sleep disturbance Average of < 7 h total time immobile per night between 8pm and 8am over one week (actigraph) plus ≥ 2 episodes of nighttime behaviors (SDI; caregiver-reported)
Stahl, 2004 [66]*	US, Europe, Canada	NR	1,698 (across three RCTs)	Mild to moderate probable or possible AD	Insomnia/sleep problems derived from physician verbatim accounts
Taameeyapradit, 2014 [67]	Thailand	Hospital	158	Probable or possible AD, mixed VaD and probable or possible AD, unspecified dementia	Sleep problems defined using a Thai version of the NPI (caregiver-reported)
Tractenberg, 2003 [4]	US	NR	104	Probable or possible AD	Sleep disturbances: < 7 h of nighttime sleep during the 2–3 weeks prior to the study and/or ≥ 2 nighttime awakenings within the previous 2 weeks (SDI, caregiver-reported; actigraphy)
Tractenberg, 2005 [68]	US	NR	662 (AD: 263, control: 399)	Probable or possible AD and non-demented elderly cohort	Sleep disturbance, defined using the SDSQ and reports of 'usual' bed and wake times
Wade, 2014 [69]*	UK/US	Outpatients	80	Mild to moderate probable or possible AD	Insomnia, defined as PSQI ≥ 6 (PSQI completed by investigator with the caregiver or patient; caregiver report prioritized)
Yin, 2015 [70]*	China	Clinic/hospital	156	Mild to moderate probable or possible AD	Sleep disturbances defined as > 2 awakenings during the night (caregiver-reported) and assessed and confirmed by PSG
Zhou, 2019 [71]	China	Neurology clinic	176 (AD = 84, control = 92)	Probable or possible AD and negative controls	Sleep disturbances defined using the PSQI

AD, Alzheimer's disease; AIS, Athens Insomnia Scale; BEHAVE-AD, Behavioral Pathology in Alzheimer's Disease; CUSPAD, Columbia University Scale for Psychopathology in Alzheimer's Disease; DLB, Dementia with Lewy bodies; DSM-5, Diagnostic and Statistical Manual of Mental Disorders, Fifth Edition; ESS, Epworth Sleepiness Scale; ICD, International Classification of Diseases; MCI, Mild Cognitive Impairment; MOUSEPAD, Manchester and Oxford Scale for Psychopathological Assessment in Dementia; NPI, Neuropsychiatric Inventory; NR, not reported; PSG, polysomnography; PSQI, Pittsburgh Sleep Quality Index; SDI, Sleep Disorders Inventory; SDSQ, sleep diary-derived sleep quality measure; UK, United Kingdom; US, United States; VaD, vascular dementia.

Table 2
Studies reporting association between sleep disturbances and clinical burden in AD patients

Author	Relevant objectives assessed	Clinical outcome results
Bliwise, 1995 [19]	The impact of the severity of AD on behaviorally defined sleep disturbances	Greater functional impairment (ADL) was associated with moderately disturbed sleep; no significant correlation was observed between severe sleep disturbances and functional impairment (ADL).
Ownby, 2014 [58]	Subtypes of sleep disturbance in patients with AD, and the relation to patient characteristics	Both the moderate and severe sleep groups had worse cognitive function (MMSE), lower functional status (BDRC), and higher depression scores (CSDD) versus the normal sleep group.
McCurry, 1999 [47]	Frequency, predictors and impact of sleep problems in AD patients	Numerically greater memory (RMBPC) impairment, humanistic burden (BDRS) scores and depression (RMBPC) with increased frequency of sleep disturbances (defined by caregivers). Prevalence OR: 1.6 for 3-point functional status (BDRS) score change in patients with sleep disturbance in past week.
Tractenberg, 2003 [4]	Efficacy of the Sleep Disorders Inventory (SDI) for assessing symptoms of sleep disturbance/disorder	No significant difference in functional impairment (ADL) for short sleep duration group versus ≥ 6 h TST; 39.1 versus 42.9. SDI scores were significantly associated with functional impairment (ADL) scores.
Tractenberg, 2005 [68]	Prevalence rates of sleep disturbance symptoms in those with possible AD, and a normal elderly control group, and the association of cognitive status with greater prevalence, worse symptomatology, or a different range of symptoms.	Greater functional impairment (ADL and IADL) for short sleep duration group versus > 6 h TST; 0.88 versus 2.60 and 1.9 versus 7.0, respectively. Higher functional impairment (ADL) scores (less impairment) for sleep problems group versus no sleep problems; 3.5 versus 2.2. No significant difference in functional impairment (IADL) for sleep problems group versus no sleep problems; 7.6 versus 6.6.
McCurry, 2006 [51]	Impact of higher % sleep on (caregivers' reports of sleep disturbances in persons with AD and actigraph records of patients' sleep-wake activity) on clinical outcomes.	Greater functioning (less functional impairment [IADL]) in those with higher % sleep Less daytime sleepiness (ESS) in those with higher % sleep Better QoL (QoL-AD) in those with higher % sleep Better physical QoL (SF-36) in those with higher % sleep
Garcia-Alberca, 2013 [29]	The association between sleep disturbances and neuropsychiatric symptoms, cognitive and functional status of patients, and severity and duration of dementia	Worse functioning (B-ADL), worse cognition (MMSE, RAVLT-IR, RAVLT-DR and TMT-B), depression (NPI) and aberrant motor behavior (NPI) were associated with sleep disturbances
de Oliveira, 2014 [25]	Factors related to sleep satisfaction in AD patients	No correlation between cognition (MMSE and clock drawing test) and sleep satisfaction. No difference in functional impairment (ADL or IADL) for unsatisfied sleep group versus satisfied sleep (4.97 versus 5.02 and 15.32 versus 13.97, respectively). Greater behavioral disturbances (NPI total score) and higher dysphoria, anxiety and apathy scores (NPI) in patients with unsatisfied sleep versus satisfied sleep. Significance for a decrease of 0.079 h of sleep for each point increase in anxiety score and for an increase of 0.075 h sleep for each point increase in apathy scores. No association between functional impairment (ADL) and sleep duration.
Ribeiro, 2018 [61]	Impact of sleep disorders, and manifestation in people with AD	Greater anxiety and depression (HADS) for sleep disturbances versus no sleep disturbances.
Yin, 2015 [70]	Five-year effect of nocturnal sleep disturbances on the long-term outcome in AD patients	No significant differences in cognition (MMSE) or functional impairment (ADL) for sleep disturbances versus no sleep disturbances groups at baseline. Significantly worse cognition (MMSE) and functional impairment (ADCS-ADL) for sleep disturbances group versus no sleep disturbances after five years. Significant increase in cumulative incidence of psychotic symptoms, eating problems and sundowning syndrome for sleep disturbances group versus no sleep disturbances after five years.

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Table 2
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Author	Relevant objectives assessed	Clinical outcome results
Leng, 2020 [44]	Sleep quality and health-related QoL in older adults with subjective cognitive decline, mild cognitive impairment, and AD	Total (SF-36), physical (PHCS SF-36), and mental (MHCS SF-36) QoL were associated with subjective sleep quality; moderate correlations between HRQoL and sleep quality. However, the AD group showed a weaker correlation than the other patient groups. Total and physical QoL were also moderately associated with sleep disturbances and total QoL alone was associated with sleep duration.
Zhou, 2019 [71]	Characteristics of sleep status and BPSD among AD patients in Eastern China, and the relationship among sleep disorder, behavioral and psychological symptoms of dementia, and cognition	Negative correlation between PSQI scores and cognition scores (MMSE) and functional impairment scores (ADCS-ADL). PSQI was closely related to behavior/neuropsychological impairment scores (NPI total score) and depression and apathy scores (both NPI).
Lebrija, 2016 [41]	Sleep disorders and neuropsychiatric symptoms in Mexican patients with AD	Significant association between anxiety (NPI) and difficulty falling asleep (versus patients without difficulty to initiate sleep), waking up at night (versus patients not waking up at night to start the day) and early morning awakenings (versus patients without early morning awakening). Significant association between aberrant motor behavior (NPI) and waking up at night (versus patients not waking up at night), wandering at night (versus patients not wandering at night), night awakenings (versus patients without night awakenings) and early morning awakenings (versus patients without early morning awakening). Significant association between agitation (NPI) and waking up at night to start the day (versus patients not waking up at night to start the day).
Kabeshita, 2017 [37]	Sleep disturbances and other behavioral and psychological symptoms of dementia at different stages of AD	Behavior/neuropsychological impairment (NPI total score) was associated with sleep disturbances in patients with low CDR; no correlation in more severe AD
Chew, 2019 [23]	Sleep duration and progression of cognitive decline in subjects with mild cognitive impairment and mild AD	Longer sleep duration in patients with rapid cognitive decline (CDR) versus those without
Escudero, 2019 [28]	Association between neuropsychiatric symptoms and cognitive and functional decline in frontotemporal degeneration and AD	Sleep problems were predictive of cognitive decline after 2.5 years based on MoCA, but sleep problems at baseline were not associated with cognitive decline after 2.5 years based on MMSE. The model on Δ Lawton (functional impairment – Lawton scale) reached significant values showing that variability in Δ Lawton was explained by the scores for depression and for sleep problems.
Mulin, 2011 [55]	Apathy and sleep/wake patterns in individuals with AD using ambulatory actigraphy	Greater WASO, time in bed, daytime inactivity periods and WASO normalized for time in bed in patients with apathy versus patients without apathy.
Moran, 2005 [54]	Sleep disturbance and its clinical correlates in a memory clinic population of AD patients	No significant difference in cognitive decline (MMSE, CAMCOG) and functional impairment (IADL, BDRS) between patients with and without sleep disturbances. Significantly more aggressiveness (BEHAVE-AD) and worse behavioral disturbances (Global rating [BEHAVE-AD]) in patients with versus without sleep disturbances.
Leger, 2017 [43]	Impact of apathy or severity scores in patients with AD on TST at night	Negative correlation between TST and ADCS-ADL score, meaning TST was positively associated with functional disability. Increased TST (AES-I), decreased awakenings (AES-I), and increased sleep efficiency (AES-I) in patients with apathy versus patients without.

(Continued)

Table 2
(Continued)

Author	Relevant objectives assessed	Clinical outcome results
Shin, 2014 [63]	Impact of nighttime sleep on cognition and behavioral and psychological symptoms of dementia in AD	No significant correlation between PSQI total scores and cognition (MMSE [Korean version]) or functional impairment (B-ADL, SI-ADL) in AD group. Apathy/indifference (NPI [Korean version]) significantly associated with PSQI total scores. Sleep latency was negatively associated between and visuospatial functions and praxis; sleep efficiency was positively associated with praxis, K-BNT and RCFT; sleep duration influenced praxis. After adjusting for age and education sleep latency was significantly associated with praxis, immediate recall and recognition, and sleep duration and efficiency correlated with praxis.

AD, Alzheimer's disease; ADCS-ADL, Alzheimer's Disease Cooperative Study-Activities of Daily Living; ADL, Activities of Daily Living; AES-I, Informant-rated Apathy Evaluation Scale; B-ADL, Bayer Activities of Daily Living; Ba-ADL, The Barthel Activities of Daily Living Index; BDRS, Blessed Dementia Rating Scale; BEHAVE-AD, Behavioral Pathology in Alzheimer's Disease; CAMCOG, Cambridge Examination for Mental Disorders of the Elderly; CDR, Clinical Dementia Rating; CSDD, Cornell Scale for Depression in Dementia; ESS, Epworth Sleepiness Scale; HADS, Hospital Anxiety and Depression Scale; IADL, Instrumental Activities of Daily Living; K-BNT, Korean version of the Boston Naming Test; MCI, mild cognitive impairment; MHCS, Mental Health Component Score; MMSE, Mini-Mental State Examination; MoCA, Montreal Cognitive Assessment; NPI, Neuropsychiatric Inventory; NR, not reported; PHCS, Physical Health Component Score; QoL, quality of life; QoL-AD, Quality of Life in Alzheimer's Disease; RAVLT-IR, Rey Auditory Verbal Learning Test; RCFT, Rey-Osterrieth Complex Figure Test; RMBPC, Revised Memory and Behavior Problems Checklist; SI-ADL, Seoul-Instrumental Activities of Daily Living Scale; SNSB, Seoul Neuropsychological Screening Battery; STAI Y-1, State Trait Anxiety Inventory for Adults; TMT-B, Trail-making test; TST, total sleep time; WASO, wake after sleep onset.

AD patients with and without sleep disturbances, but behavioral disturbances were significantly associated with sleep disturbance [54]. Two studies reported that longer sleep duration was associated with cognitive/functional impairment [23, 43]. In general, however, most studies have reported that poor sleep impacts both cognition and functional impairment.

Behavioral symptoms and neuropsychiatric functioning of AD patients are also worse among those with sleep disturbances, [25, 29, 41, 47, 54, 71] particularly depression and anxiety [25, 29, 58, 61, 71]. Several studies also reported significant associations between poor sleep and apathy in AD patients [55, 71]. Mulin et al. found that those with AD and apathy had reduced daytime activity levels and poorer nocturnal sleep quality [55]. Increased sleep and decreased circadian stability have also been associated with apathy in AD patients [25, 43]. Moran et al. found that sleep disturbances were associated with aggressiveness and greater global ratings for behavioral problems in AD patients [54]. Other studies reporting clinical impacts of sleep disturbances are summarized in Table 2.

Sleep problems are also associated with reduced QoL in AD patients [44, 47, 51]. Leng et al. reported that sleep disturbances, poor subjective sleep quality, and shorter sleep duration were all associated with reduced mental, physical or total QoL scores [44]. In line with this, greater actigraphically-defined sleep

efficiency has been associated with better physical QoL scores in AD patients [51].

Longitudinal studies suggest that sleep disturbances in AD may predict worsening clinical symptoms. Yin et al. found significantly lower MMSE and Alzheimer's Disease Cooperative Study-Activities of Daily Living (ADCS-ADL) scores after five years in those reporting sleep disturbances, despite comparable scores in good versus poor sleepers at baseline [70]. They also reported a higher cumulative incidence of psychotic symptoms and sundowning syndrome among those with sleep disturbances after five years [70]. Escudero et al. also reported that sleep problems were predictive of cognitive and functional decline in AD patients (Montreal Cognitive Assessment [MoCA]) [28].

Caregiver burden and institutionalization of patients

Impact of insomnia/sleep disturbances on caregiver burden

Caregiver burden associated with sleep problems in AD patients was overall found to be increased (20 of 21 studies; Table 3). Only one study reporting caregiver burden outcomes included patients with a sleep disturbance defined as insomnia (30.4% with clinically diagnosed insomnia) [56]. Caregivers were typically spouses or adult children/grandchildren,

Table 3
Studies reporting on the association between patient with AD and sleep disturbances and caregiver burden

Author	Study objective	Caregiver characteristics			Clinical outcome result
		Relationship to patient	Age (y)	Gender	
Owby, 2010 [57]	Ethnic differences in sleep problems among patients with AD	Spouse: 44.05% Child/grandchild: 48.39% Other relative: 5.32% Other: 2.02%	NR	F: 105 M: 286	Positive association (RC) between patients' difficulty falling asleep, nighttime awakenings, and patient early morning awakenings and caregiver depression (CES-D).
Owby, 2014 [58]	Subtypes of sleep disturbance in patients with AD, and the relation to patient characteristics and caregiver mood	Most frequently spouse or adult child; study does not report exact figures	Mean (SD): 57.3 (14.9)	F: 316 M: 134	Higher caregiver depression (CES-D) scores for caregivers of the moderate and severe sleep disturbances groups versus normal sleep group.
McCurry, 2004 [49]	Anxiety and nighttime behavioral disturbance in a community-dwelling sample of patients with AD	Cohabiting with patient: 94% See patient everyday: 6% (female spouse: 70%)	Mean (SD): 70.1 (12.8)	Female spouses: 70%	Anxiety doubled the odds of awakening the caregiver at night (caregiver reported) (OR 2.0, 95% CI: 1.4, 2.9). Increased patient functional impairment in activities of daily life was significantly related to awakening caregiver at night (caregiver reported) (OR: 1.6, 95% CI: 1.2, 2.3). No significant relationship between patient depression and awakening caregiver at night (caregiver reported).
McCurry, 2005 [50]	Impact of comprehensive sleep education program NITE-AD on sleep in dementia patients living at home with their family caregivers	Spouses: 58%	21–87	F: 72%	92% of caregivers of patients with sleep problems reported poor sleep themselves (PSQI) (baseline).
McCurry, 2006 [51]	Caregivers' reports of sleep disturbances in persons with AD and actigraphic records of patients' sleep-wake activity, and discrepancies in this relationship	Spouses: 54% Remaining were adult relatives living with patient	21–87	F: 65% M: 35%	Greater caregiver misperception of patient poor sleep associated with higher caregiver burden (SCB) and more complaints of caregiver poor sleep (PSQI, CSQ).
McCurry, 1999 [47]	Frequency, predictors, and impact of sleep problems in a population-based sample of 205 AD patients	Spouse: 73% Adult child: 6% Other: 7% Lived away from patient but saw them at least weekly: 14%	32–94 Mean: 68	F: 65%	Positive correlation between increased frequency of all sleep disturbances (except nightmares) and distress (RMBPC). 70% and 69% of caregivers reported 'awakening caregiver at night' and 'patient sleeping less than usual' as moderately-severely distressing (RMBPC), respectively.
Gehrman, 2018 [31]	Association between symptoms of disturbed sleep in patients with AD and caregiver burden outcomes.	Spouse: 81% Child: 15% Other 4%	Mean (SD) 66.1 (11.7)	F: 61% M: 39%	Significant associations between nocturnal awakenings, nocturnal wanderings and daytime sleepiness compared with caregiver burden (SCB)
Lukovits, 1992 [45]	To compare family members' and nursing staff members' perceptions regarding behavioral disturbances in patients with severe AD	Nurses and family members	NR	NR	Nurses reported more concern (5-point rating scale) than family for vegetative behaviors (including sleep disturbances); concern decreased with AD severity.

(Continued)

Table 3
(Continued)

Author	Study objective	Caregiver characteristics			Clinical outcome result
		Relationship to patient	Age (y)	Gender	
Grace, 2000 [32]	Frequency of sleep disturbances in patients with AD and DLB and impact on caregivers	NR	NR	NR	Sleep disturbances were moderately-very severely distressing (NPI) in 8/20 caregivers.
Donaldson, 1998 [26]	Impact of subgroups and individual symptoms of non-cognitive disturbance on the carers of AD patients	Lived with patient: 74% Spouse: 53% Child: 36% Other relatives/ friends: 11%	32–87 Mean (SD): 63.1 (13.6)	F: 57%	Positive correlations between sleep disturbances and caregiver burden (GSS) and distress (GHQ); sleep disturbances predicted distress in caregivers.
de Oliveira 2014 [25]	Sleep satisfaction in patients with AD	NR	NR	NR	No significant difference in caregiver burden (ZBI Brazilian version) for unsatisfied sleep versus satisfied sleep.
Camargos, 2015 [22]	Trazodone on cognitive performance in patients with AD	NR	NR	NR	70.7% of caregivers of patients with sleep disturbances reported moderate/extreme distress (NPI); 7.3% reported mild distress.
Kazui, 2016 [38]	Differences of trajectories of 12 kinds of BPSDs by disease severity in four major dementias showing the frequency, severity, and associated caregiver distress of BPSDs using (J-BIRD)	NR	Mean (SD) 76.9 (8.7)	F: 752 M: 339	21.2%, 21.4%, 43.4%, and 76.5% of caregivers of patients with sleep disorders and CDR 0.5, CDR 1, CDR 2 and CDR3 had moderate or greater distress (NPI-D), respectively. Significant differences between associated caregiver distress (NPI-D) between CDR groups 0.5–3.
Okuda, 2019 [56]	Sleep disturbance in AD patients and the burden on, and health status of, their caregivers in Japan	Spouse: 4.8% Child/grandchild (including son/daughter-in law): 88.7% Other: 6.5%	Mean (SD): 50.4 (12.4)	F: 50.2% M: 49.8%	Significant positive correlation between patient sleep disturbances and total care burden (BIC-11). Significant positive association between patient sleep disturbances and caregiver time-dependent emotional, existential, physical and service-related, and total burden (BIC-11), caregiver depression (Japanese PHQ-9) and caregiver poor sleep quality (Japanese SF-12v2). Significant negative association between patient sleep disturbances and caregiver physical and mental QoL (Japanese SF-12v2).
Allegri, 2006 [15]	Predictive value of behavior-related burden on AD caregivers	Spouse: 54.6% Child: 36% Other: 9.4%	Mean (SD): 59.6 (14.8)	F: 67 M: 15	Positive correlation between patient sleep disturbances and patient abnormal behavior (wandering) compared with caregiver burden/distress (ZBI).
Yin, 2015 [70]	5-year effect of nocturnal sleep disturbances on the long-term outcome in AD patients	Spouse: 100%	NR	NR	Increased caregiver negative emotions in group caring for patients with sleep disturbances versus those caring for patients with no sleep disturbances after 5 years.

Taameeyapradit, 2014 [67]	Characteristics of the BPSD and its severity among patients with dementia and their caregivers' stress	NR	20–39: <i>n</i> = 30 (18.99%) 40–59: <i>n</i> = 109 (68.98%) 60+: <i>n</i> = 19 (12.02%)	F: 116 (73.42%) M: 42 (26.58%)	Patient sleep disturbances rated in top four most burdensome (NPI Thai variant) symptoms (higher than irritability, depression, and anxiety). 33.33% of caregivers of patients with sleep problems reported high suffering and were not capable of managing problem; 31.11% reported they were greatly troubled and patient sleep problems were difficult to manage; 20.00% reported considerable trouble and not easily managed.
Pang, 2002 [59]	Cross-regional and cross-cultural differences in symptom-related caregiver distress due to behavioral problems of patients with AD	Spouse: 37.9–61.8% Child/sibling/child's spouse: 36.0–61.3% Friends/other: 0–15.5%	Mean (SD): US: 60 (14) Taiwan: 59 (16) Hong Kong: 58 (14)	F:M ratio: US: 1.7:1 Taiwan: 2:1 Hong Kong: 1:1	Patient sleep problems were moderately-very severely distressing (NPI) in ~35% of caregivers from all ethnic groups (Taiwanese, Hong Kong Chinese and Caucasian American).
Hart, 2003 [33]	Behavioral and psychological symptoms in patients with a diagnosis of established AD for at least three years	Carer or next-of-kin	NR	NR	Patient sleep disturbances were one of the least well tolerated BPSD (alongside aggression and irritability), causing high caregiver distress (NPI), despite medications*
Tractenberg 2003 [4]	Efficacy of the SDI as an instrument for assessing symptoms of sleep disturbance/ disorder	Live in caregivers (residing with patient)	47–92	F: 49%	Highest caregiver distress (NPI) for nighttime wandering and awakening caregiver. Lower caregiver distress related to SDI items in patients with ≥ 6 h TST versus < 6 h TST.
Kim, 2017 [39]	Impact of sleep disturbance in caregivers of AD	NR	NR	NR	Significant positive correlation between patient sleep problems and caregiver distress (NPI).

*High observed rates of drug use including cholinesterase inhibitors, antipsychotics, and antidepressants among patients. For studies using the NPI to assess caregiver distress, only the sleep disturbance item was considered for this review. AChEI, acetylcholinesterase inhibitor; AD, Alzheimer's disease; BIC-11, Burden Index of Caregivers-11; BPSD, behavioral and psychological symptoms of dementia; CDR, Clinical Dementia Rating; CES-D, Center for Epidemiological Studies Depression Scale; CSQ, Caregiver Sleep Questionnaire; DLB, Dementia with Lewy bodies; F, female; FTL, frontotemporal lobar degeneration; GHQ, General Health Questionnaire; GSS, Gileard's Strain Scale; HAMA, Hamilton Anxiety Scale; HDRS, Hamilton Depression Rating Scale; M, male; NITE-AD, Nighttime Insomnia Treatment and Education for Alzheimer's Disease; NPI, Neuropsychiatric Inventory; NR, not reported; n.s, not significant; PHQ-9, Patient Health Questionnaire-9; PSS, Perceived Stress Scale; PSQI, Pittsburgh Sleep Quality Index; QoL, quality of life; RC, regression model coefficient; RCT, randomized controlled trial; RMBPC, Revised Memory and Behavior Problem Checklist; SCB, Screen for Caregiver Burden; SD, standard deviation; SDI, Sleep Disorders Inventory; SF-12v2, 12-item Short Form Health Survey version; SF-36, 36-Item Short Form Health Survey; VaD, vascular dementia; ZBI, Zarit's Burden Interview.

and frequently aged over 50 years, with a mean reported caregiver age of 60–80 years. Ownby et al. found sleep disturbances in AD patients were associated with greater caregiver depression, but cognitive decline was not controlled for in the study [58]. To this point, Kazui et al. reported a significant increase in sleep disturbance-associated distress among caregivers of patients with more severe AD [38]. The type of sleep problem is also a factor; McCurry et al. reported that nighttime awakenings, but not excessive sleeping, were most disturbing to caregivers and associated with greater memory impairment and worse functional status in patients [47].

The literature overwhelmingly suggested that sleep disturbances and disruptive nighttime behaviors (awakenings and wanderings) in AD patients are associated with considerable distress, poor health status and QoL, and reduced sleep in caregivers [4, 15, 22, 26, 31, 33, 38, 39, 47, 50, 56, 59, 67]. Interestingly, objective caregiver burden has also been associated with mis-perceived poor sleep quality in patients by caregivers [51]. Only one identified study reported that sleep disturbances were not associated with caregiver burden; de Oliveira et al. reported no significant difference between the burden in caregivers of patients with satisfactory versus unsatisfactory sleep [25].

Sleep disturbances in AD patients are also associated with caregiver depression, anxiety and mood, [51, 56–58, 70] and have a negative impact on caregivers' overall QoL [31, 56]. Caregivers' sleep worsened in the presence of patients' sleep disturbances; [51, 56]. Okuda et al. found sleep disturbances in AD patients correlated significantly with higher caregiver PSQI scores (worse sleep quality) [56].

Several studies investigated the impact of specific sleep characteristics and behaviors on caregiver burden. Nighttime behaviors (awakenings and wanderings) had the largest impact on caregiver burden (four studies) [4, 15, 31, 47]. Notably, those with anxiety were twice as likely to waken their caregiver versus those without, and heightened anxiety was reported among patients taking sedative or hypnotic, antianxiety, antipsychotic or antidepressant medications, suggesting that the presence of anxiety led to the prescription of these medications [49].

Institutionalization of AD patients with sleep disturbances

Insomnia (two studies) [17, 18] and sleep disturbances (one study) [70] are key factors influencing the institutionalization of AD patients. In a five-year

study, more AD patients with sleep disturbances (caregiver report of ≥ 2 nighttime awakenings) lived in special dementia care units than those without sleep disturbances [70]. Baek et al. reported that a greater proportion of AD patients with insomnia were admitted to a long-term care facility and had longer durations of admission than those without insomnia [17]. Insomnia was also independently associated with admission to a nursing home after discharge from a dementia unit [18]. Importantly, other factors influence institutionalization, such as the availability of home services [18].

Thus insomnia/sleep disturbances may both accelerate the rate of deterioration of AD patients and lead to institutionalization. Interestingly, a 2006 study by McCurry et al. reported that misperception of patient sleep by caregivers in comparison to actigraphy was associated with greater objective caregiver burden [51]. Caregivers play a key role in the reporting and management of sleep disturbances in AD patients, and their perception of patients' sleep disturbance likely influences patient management, including the decision to admit AD patients into care facilities.

Treatment of sleep disturbances in Alzheimer's disease

Few insomnia treatment studies have been conducted in AD patients. We identified only 23 interventional studies and the majority were small (14 studies with ≤ 50 subjects), highlighting the need for further studies to assess treatment efficacy and patient and caregiver outcomes in this population.

Non-pharmacological treatments

Of eight studies investigating non-pharmacological treatments, some found improvements not only in patients' sleep, but also other patient and caregiver outcomes (Table 4). Lee et al. reported improvement in subjective, but not objective, sleep (PSQI) in AD patients with insomnia treated with timed blue light, as well as improvements in behavioral disturbances and caregiver distress [42]. Ancoli-Israel et al. reported that daytime bright light exposure increased nocturnal sleep consolidation [16]. In contrast, some studies did not find light therapy effective in improving sleep overall [24, 27]. Of these studies, however, Dowling et al. found that people with the most impaired rest-activity rhythm responded to light therapy, [27] and van Someren et al.'s 1999 reanalysis of Colenda et al.'s 1997 study found that light therapy improved interdaily rhythm stability [24, 73].

Table 4
Impact of non-pharmacological treatments on sleep and clinical outcomes in AD patients

Author	Study objective	Intervention and comparator	Clinical outcome result
Ancoli-Israel, 2003 [16]	Impact of light on sleep and circadian activity rhythms in patients with probable or possible AD	Morning bright light (2,500 lux) versus morning dim red light (<300 lux) versus evening bright light (2,500 lux) (average of 92.1 min treatment per 120 min session)	No light treatment resulted in significant effects on actigraphically-measured sleep during night or day, using traditional sleep measures; TST, WASO, % sleep, % wake, number of nighttime awakenings, average length of nighttime awakenings, number of daytime naps, length of time between naps (wrist actigraphy). Bouts of sleep at night (computed using actigraphy data) increased in the morning bright light group and evening bright light group at day 6–10; no significant change with dim light. No significant difference in bouts of wake during the day (computed using actigraphy data) versus baseline for any group at day 6–10. Mean wake-bout length decreased from day 6–10 to post-treatment follow up in the evening bright light group (rebound/relapse effect). Activity rhythms (measured by wrist actigraphy) improved with evening bright light only for mean and maximum activity levels.
Colenda, 1997 [24]	Effects of phototherapy delivered by light visors on disturbed sleep patterns of community-dwelling research subjects with AD	10 consecutive days of 2,000 lux of full spectrum bright light (2 h each morning)	No significant changes in circadian rest-activity (actigraphy) at post-treatment versus baseline, except one subject. van Someren et al.'s 1999 reanalysis found that a nonparametric procedure indicated that light therapy improved interdaily rhythm stability [73]. Significant increase in TST (actigraphy) at post-treatment versus baseline for one subject and significant decrease for one subject. No changes in nighttime awakenings (actigraphy) with treatment.
Dowling, 2005 [27]	Effectiveness of morning bright light therapy in reducing rest-activity (circadian) disruption in institutionalized patients with severe AD	Morning bright light (≥ 2500 lux) versus usual room light (control) (treatment for 5 days a week for 10 weeks)	No significant changes in sleep efficacy, nighttime sleep time, nighttime wake time, number of nighttime awakenings, daytime wake time (all wrist actigraphy) with treatment versus control and no overall improvement in circadian rhythms (wrist actigraphy: parametric and nonparametric analysis) with treatment versus control; only subjects with the most impaired rest-activity rhythm responded to the light therapy.
Lee, 2018 [42]	Effect of timed blue-enriched light on subjective and objective sleep in AD patients and to evaluate changes in their caregiver burden	Timed blue enriched light (1 h per day for two weeks) and control group	Subjective sleep (PSQI) scores significantly decreased with treatment/ no significant changes in objective sleep with treatment versus baseline. Patient's neuropsychiatric symptoms (NPI-S) and caregiver distress (NPI-D) scores reduced after treatment; not significant.
McCurry, 2005 [50]	NITE-AD in dementia patients living at home with their family caregivers.	NITE-AD (combination of sleep hygiene, daily walking, and light exposure intervention over three weekly treatment sessions) versus control (general dementia education and caregiver support)	Improvement in time awake at night (wrist actigraphy) from baseline with treatment; maintained over 6-month follow up. Fewer nighttime awakenings (wrist actigraphy) with treatment versus control (32% reduction from baseline). Significantly lower levels of depression (RMBPC) with treatment versus control, maintained over 6-month follow up. Lower daytime sleepiness (ESS) with treatment versus control over 6-month follow up (controlling for MMSE).

(Continued)

Table 4
(Continued)

Author	Study objective	Intervention and comparator	Clinical outcome result
McCurry, 2011* [52]	Effects of walking, light exposure, and a combination intervention (walking plus light plus sleep education) on the sleep of persons with AD	NITE-AD versus walking versus light therapy (~2500 lux) versus control	Reduced total wake time at night (wrist actigraphy) for walking, light and NITE-AD with treatment versus control. No difference between active treatment groups. Improvements not maintained over time in 6-month longitudinal analysis. Trend for fewer awakenings (wrist actigraphy) for the walking group versus control. Moderate effect size improvements in sleep percent (wrist actigraphy) for active treatment groups. No significant difference in TST (wrist actigraphy) and caregiver-reported patient sleep (SDI) between groups.
McCurry, 2003 [48]	Feasibility of training caregivers to implement sleep hygiene recommendations in dementia patients	Caregivers received training and tailored recommendations/education to aid their implementation of sleep hygiene strategies among patients versus nondirective support	Patients adhered to sleep hygiene goals (consistent bedtime, consistent rising time, nap restriction, walking) significantly more consistently in active training group. Changes in sleep hygiene behaviors occurred more often if caregivers received specific suggestions/assistance
Simoncini, 2015 [64]	Effectiveness of acupressure for the treatment of insomnia and other sleep disturbances, and to show that the acupressure treatment is feasible also in elderly resident patients	Daily acupressure for 8 weeks	Positive subjective perception of sleep after treatment and maintained at follow up; hours of sleep perceived to be increased; time to fall asleep decreased; quality of sleep increased. Significant improvement in general health (GHQ28) with treatment versus baseline: PSQI significantly correlated to GHQ28 with good sleep encouraging perception of better health. Positive difference found with regard to sleep and mood. No change in cognitive functioning (MMSE) and functional status (ADL, IADL) with treatment versus baseline. Improvement in behavioral and psychological functioning (NPI) and positive non-significant trend of improvement in anxiety (STAI Y-1) after treatment.

*Post-test outcomes controlled for baseline age, gender, depression, comorbidity limitations, MMSE and sleep apnea scores. ADL, Activities of Daily Living; ESS, Epworth Sleepiness Scale; GDS, Global Deterioration Scale; GHQ28, Global Health Quality of Life; IADL, Instrumental Activities of Daily Living; MMSE, Mini-Mental State Examination; NITE-AD, Nighttime Insomnia Treatment and Education for Alzheimer's Disease; NPI, Neuropsychiatric inventory; PSQI, Pittsburgh Sleep Quality Index; STAI Y-1, State Trait-Anxiety Inventory; TST, total sleep time; WASO, wake after sleep onset.

Acupressure treatment was assessed in one study and improved sleep in elderly institutionalized AD patients with insomnia, but had no effect on cognition (MMSE) or functional status (ADL) [64].

McCurry et al. reported that an approach combining sleep hygiene, walking, and light exposure (NITE-AD) reduced the frequency and duration of nighttime awakenings, and lowered daytime sleepiness (Epworth Sleepiness Scale [ESS]) and levels of patient depression (RMBPC) versus controls [50]. They also reported improvements in patients' actigraphically-measured total wake time with NITE-AD versus controls; similar results were reported

for the groups receiving light and walking therapies [52]. However, caregiver-reported subjective outcome measures (Sleep Disorders Inventory [SDI]) did not replicate these results, but caregiver reports are not always consistent with actigraphy [52]. Unfortunately, improvements in sleep with any interventions were not sustained at six months, likely due to reduced adherence over time [52]. Indeed, in the latter NITE-AD study, patients adhering to recommendations ≥ 4 days/week had significantly better sleep outcomes [52]. These findings highlight a key challenge with non-pharmacological treatment, which may not always be feasible in patients with AD and

other comorbidities due to inability or lack of willingness to partake. Notably, active training and support for caregivers led to a greater success in implementing good sleep hygiene practices in AD patients [48].

Pharmacological treatments

Only 15 studies investigated the effects of pharmacological sleep treatments in AD patients (Table 5). Although several treatments showed promise in improving clinical and caregiver symptoms, inconsistent clinical results were also apparent and there are insufficient data supporting many commonly used treatments.

Benzodiazepines and non-benzodiazepine 'z-drugs'

A small study ($N=7$) by McCarten et al. investigated the efficacy of the benzodiazepine triazolam (0.125 mg at bedtime) in patients with AD but failed to detect significant effects on sleep parameters versus placebo (Table 5) [46]. Yin et al. reported improvements in sleep quality (PSQI) and daytime sleepiness (ESS) with the non-benzodiazepine zolpidem tartrate, versus no treatment in a 5-year study; no change was seen in MMSE, ADCS-ADL or NPI scores with zolpidem tartrate after five years versus no treatment, suggesting no significant impact on cognitive decline or neuropsychiatric behaviors, but no safety outcomes were reported [70].

Melatonin/melatonin receptor agonists Two studies found melatonin to be ineffective at improving sleep, agitation scores, or circadian rhythm parameters in AD patients based on actigraphy [30, 65]. Conversely, Brusco et al., reported significant improvement in sleep quality and decreased sundowning with melatonin 9 mg, but this was an open label study with only 14 patients [20]. Similarly, Wade et al. reported significantly improved sleep efficiency among AD patients with insomnia taking prolonged-release melatonin in a double-blind, placebo-controlled study after 24 weeks, but PSQI scores were not significantly different between groups [69]. Cognition was also improved with melatonin versus placebo (for MMSE and AD Assessment Scale-Cognition scores), as well as self-care and activities of daily living scores (IADL) [69].

Antidepressants A small study by Camargos et al. found that trazodone 50 mg increased sleep duration and efficiency in AD patients versus placebo, was well-tolerated, and did not increase daytime sleepiness [21]. Petrescu et al. also reported that patients

taking trazodone (50–200 mg) had more total sleep time in both patient- and nurse-reported measures, although it was associated with more side effects than quetiapine [60]. Interestingly, trazodone may have a possible neuroprotective effect in AD patients with insomnia; a retrospective study by La et al. reported that among a group of older adults, including some with AD, those not using trazodone declined 2.4-fold faster than trazodone users (median dose 50 mg) across an average of 3.75 years based on MMSE results [40]. Notably, while trazodone is an antidepressant, a dose of 50 mg is consistent with use for insomnia, rather than depressive disorders. A small study assessing the antidepressant mirtazapine reported no effects on sleep outcomes in AD patients (Table 5) [62].

Antipsychotics A five-year study by Yin et al. reported improved daytime (ESS) and nighttime (PSQI) sleep symptoms with low-dose risperidone versus zolpidem tartrate, melatonin, or no drug treatment in AD patients with sleep disturbances [70]. Risperidone also alleviated neuropsychiatric symptoms (NPI) versus no treatment and did not significantly affect cognition or functional autonomy versus zolpidem, melatonin, or no treatment, suggesting no acceleration in mental deterioration [70]. Risperidone was also associated with a significant improvement in caregiver mood and sleep and a lower expectation that a patient would be institutionalized versus the melatonin and no treatment groups, implying improved outlook; institutionalization of risperidone-treated patients was significantly lower than in patients not receiving sleep medication [70]. However, this was not a randomized trial, and treatment group assignment was based on the preference of the spousal caregiver. In a nursing home-based study, Meguro et al. reported that low-dose risperidone was effective at increasing patient sleep and decreasing daytime wandering [53]. Importantly, patients did not exhibit daytime oversleeping or side effects which would have prevented them participating in daily activities [53].

Orexin receptor antagonists Orexin receptor antagonists have emerged as a new generation of insomnia treatments. In a polysomnography study of AD patients with insomnia, Herring et al. reported that suvorexant significantly increased total sleep time versus placebo, and decreased waking after sleep

Table 5
Pharmacological treatments: impact on sleep and clinical outcomes in AD patients

Author	Study objective	Intervention and comparator	Clinical outcome result
Brusco, 1999 [20]	Melatonin in selected populations of sleep-disturbed patients	Melatonin 9 mg, for 22 to 35 months	Sleep quality (clinical interviews and sleep logs; caregiver-reported) improved and sundowning measure was no longer detected in 12/14 patients from baseline with melatonin. No differences in the evolution of disease with melatonin versus baseline based on neuropsychological/cognitive functioning tests (FAST; MMSE)
Camargos, 2014 [21]	Trazodone to treat sleep disturbances in patients with AD	Trazodone 50 mg versus placebo for two weeks	Nighttime total sleep time (actigraphy) increased by 8.5 percentage points post-treatment with trazodone; 42.5 min more of sleep with trazodone versus placebo. Daytime sleepiness and naps (actigraphy) not induced in either group. Trended towards reduction in time spent awake after sleep onset and number of awakenings (actigraphy) with trazodone; not significantly different between group. No effect on cognitive functioning (MMSE) with either treatment
Gehrman, 2009 [30]	Melatonin effects on sleep and agitation in institutionalized patients with AD	Melatonin 8.5 mg immediate-release and 1.5 mg sustained release combined dose (nightly) versus placebo for 10 days	No significant treatment effects on sleep (TST night, % sleep in night, WASO, % wake, total daytime sleep, % sleep in day, number and mean duration of sleep episodes) and circadian rhythm parameters (actigraphy) with melatonin versus placebo, and no change from treatment to follow up.
Hannesdottir, 2013 [34]	AZD5213 effects on sleep in subjects with mild cognitive impairment and mild AD	Anti-histamine AZD5213 (three different doses) versus placebo for four weeks	Dose-related increase in sleep-related AEs with two higher doses of AZD5213/ TST (PSG) reduced with two higher doses of AZD5213 versus placebo (no change with low dose). Decrease in TST did not result in impairment in next-day functioning (Psychomotor Vigilance Task and subjective reports of daytime sleepiness). Only small and non-clinically relevant effects on attention/response speed and memory accuracy (CogState tasks) with AZD5213.
Herring, 2020 [35]	Suvorexant for treating insomnia in patients with AD using sleep laboratory polysomnography experiments	Suvorexant 10 mg (increased to 20 mg based on clinical response) versus placebo four weeks	Greater LS mean change (improvement) from baseline in TST (PSG), greater proportion of patients with ≥ 50 min improvement in TST and greater LS mean change from baseline in WASO (PSG) at week 4 with suvorexant versus placebo. No differences between groups on objective cognitive (MMSE) and psychomotor tests (digit substitution), or trial partner-reported neuropsychiatric symptoms (NPI).
Ishikawa, 2016 [36]	Memantine on polysomnography variables and behavioral and psychological symptoms of dementia	Memantine 5 mg, increased by 5 mg each week up to 20 mg for four weeks	At week 4, memantine was associated with improved mean subjective sleep scores (AIS) and significant improvements in neuropsychological functioning and behaviors (NPI total score), TST (PSG), sleep efficiency (PSG), and nighttime awakenings (PSG) versus baseline. At week 4, memantine was associated with a significant decrease in anxiety (NPI) and irritability (NPI) versus baseline but did not result in a significant difference in cognitive functioning (MMSE) or dementia rating (CDR).

(Continued)

Table 5
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Author	Study objective	Intervention and comparator	Clinical outcome result
La, 2019 [40]	Long-term use of trazodone and delayed cognitive decline	Trazodone (median prescribed dose: 50 mg) versus matched non-trazodone users (over two consecutive annual visits)	Non-users' cognitive functioning (MMSE) declined 2.4-fold faster than trazodone users' across an average of 3.75 years.
McCarten, 1995 [46]	Triazolam, in patients of AD who were reported by caregivers to be frequently up at night	Triazolam 0.125 mg (two nights); patients acted as own control, receiving placebo for three nights prior- and two nights post-treatment	No significant effects on TST at night, latency to sleep onset, number of nighttime awakenings, TST in the day, mean level of activity in night or day (actigraphy) with triazolam. 3/6 with full data showed modest hypnotic response with increases in TST, but more nocturnal arousals with triazolam. Withdrawal effect reported in 3/7 patients during placebo wash-out (decreased TST at night). No significant drug versus placebo effects on memory (computerized memory test), and no relationship with patients' response to treatment.
Meguro, 2004 [53]	Risperidone in treating wandering and disturbed sleep/wake patterns in patients of AD	Risperidone 1 mg/day versus non-risperidone for one month	Daytime sleeping hours (caregiver-rated) decreased by 1.2 h, nighttime sleep hours (caregiver-rated) increased by 3.8 h and wandering hours (caregiver-rated) decreased by 2.7 h with risperidone versus pre-treatment. <i>Post hoc</i> analysis found significant difference in nighttime sleep hours between pre- and post-treatment with risperidone. No new daytime oversleeping or side effects that would have prevented participation in daily activities (caregiver-rated) with risperidone. No deterioration in cognitive functioning (MMSE and CASI) following risperidone treatment; some improvement shown for some patients (CASI).
Petrescu, 2019 [60]	Trazodone and quetiapine among inpatients clinically stable psychiatric	Trazodone 50–200 mg versus quetiapine 50–300 mg (treatment period: NR)	Longer TST (patient' subjective reports, nursing sleep logs) and daily dizziness (patients' subjective report) reported in patients taking trazodone versus quetiapine. Sleep efficiency (nursing sleep logs) similar between groups.
Scoralick, 2017 [62]	Mirtazapine in the treatment of sleep disorders in patients with AD	Mirtazapine 15 mg versus placebo for 14 days	No significant effects on nighttime TST (actigraphy) with mirtazapine versus placebo (gain of 55.3 min versus placebo). No significant decrease in WASO (actigraphy) nighttime awakenings, and % sleep with mirtazapine versus placebo. No effect on cognitive function (MMSE) or functional status (Katz scale) with mirtazapine versus placebo. Significant effect on daytime sleep duration with mirtazapine versus placebo, but no impact on naps.
Singer, 2003 [65]	2 dose formulations of melatonin for the treatment of insomnia in patients with AD	Melatonin 2.5 mg sustained release versus melatonin 10 mg immediate-release versus placebo for 8 weeks	Weak non-significant trend for more TST at night (actigraphy) in the melatonin groups versus placebo. Very weak trend for decreased day-night sleep ratio (actigraphy) and significantly greater sleep quality (caregiver-reported sleep logs) for the melatonin 2.5 mg group versus placebo. Changes in neuropsychological symptoms and behaviors (NPI) only seen in melatonin 2.5 mg group versus placebo; this was due to baseline differences and changes were not associated with changes to sleep. No significant difference in sleep-related symptoms (SDI) between groups.

(Continued)

Table 5
(Continued)

Author	Study objective	Intervention and comparator	Clinical outcome result
Stahl, 2004 [66]	Galantamine, and night time sleep related problems	Galantamine 8 mg bid versus galantamine 12 mg bid versus placebo (data from three trials; treatment period: NR)	Sleep-related AEs (physician-reported) not significantly different between groups. Significantly more overall concomitant sleep medication use in galantamine 24 mg group versus galantamine 16 mg group; no other significant differences in pairwise comparisons.
Wade, 2014 [69]	Melatonin on cognitive function and sleep in patients with mild to moderate AD	Melatonin 2 mg prolonged-release versus placebo for 24 weeks	Sleep efficiency (PSQI completed by investigator with the caregiver or patient; caregiver report prioritized), cognitive functioning (ADAS-Cog, MMSE), and functional status (IADL) improved significantly with melatonin versus placebo at 24 weeks. No difference in PSQI global scores (completed by investigator) between groups at 24 weeks. Trend for improvement in sleep quality (sleep diary) with melatonin at week 12. No differences in neuropsychological symptoms and behaviors (NPI) between groups. Decreased caregiver distress (SDI) in both groups.
Yin, 2015 [70]	5-year effect of nocturnal sleep disturbances on the long-term outcome in AD patients	Risperidone 0.5–1 mg versus zolpidem tartrate 5–10 mg versus melatonin 2.55 mg versus no drug over five years	Significantly lower institutionalization to special dementia care units and incidence of psychotic symptoms with risperidone versus no drug after 5 years. No significant differences in cumulative incidences of syndrome, cognitive functioning (MMSE), functional impairment (ADCS-ADL) and incidences of eating problems between groups after 5 years. Significantly lower psychiatric symptoms (NPI) with risperidone versus placebo. Significantly lower daytime sleepiness (ESS), significantly increased sleep quality (PSQI), caregiver depression (HAMD), less pronounced increase in caregiver anxiety (HAMA), improved caregiver sleep quality (PSQI), less caregiver daytime sleepiness (ESS) with risperidone versus other groups after 5 years; significantly lower with zolpidem tartrate versus no drug after 5 years. Significantly lower caregiver increased hope that patient would be admitted to a nursing home with risperidone versus melatonin and no drug after 5 years; no difference versus zolpidem tartrate (Caregiver emotional attitude assessed via self-made questionnaire).

AD, Alzheimer's disease; ADCS-ADL, Alzheimer's Disease Cooperative Study-Activities of Daily Living; ADAS-Cog, Alzheimer's Disease Assessment Scale – Cognition; AE, adverse event; AIS, Athens Insomnia Scale; bid: twice a day; CASI, Cognitive Abilities Screening Instrument; CDR, Clinical Dementia Rating; CGI, Clinical Global Impression; ESS, Epworth Sleepiness Scale; FAST, Functional Assessment Tool for AD; HAMA, Hamilton Anxiety Scale; HAMD, Hamilton Depressive Scale; LS, least squares; MMSE, Mini-Mental State Examination; NPI, Neuropsychiatric Inventory; NR, not reported; PSG, polysomnography; PSQI, Pittsburgh Sleep Quality Index; TST, total sleep time; WASO, wake after sleep onset.

onset, with no worsening of cognitive (MMSE) or psychomotor tests (digit symbol) [35]. Currently, suvorexant is the only pharmacotherapy specifically approved for the treatment of insomnia in AD patients, although other drugs in this class are also being tested in the AD population.

DISCUSSION AND CONCLUSION

This literature review demonstrates that sleep disturbances in AD patients are associated with greater cognitive decline, poorer functional ability and reduced QoL. Furthermore, behavioral symptoms

and neuropsychiatric functioning of AD patients, including depression, apathy, psychotic symptoms, and sundowning syndrome, are more frequent and severe among those with sleep disturbances. Sleep disturbances and associated disruptive nocturnal behaviors in AD patients also result in considerable caregiver burden, including increased distress, depression and sleep problems, and decreased QoL. Sleep disturbances are a major factor influencing the institutionalization of AD patients, likely through their impact on caregiver burden. Previous reviews have also identified nighttime awakenings and troublesome behaviors (e.g. wandering) as the most burdensome symptoms and key motivating factors for institutionalization [1, 74].

In addition, sleep problems have been associated with increased risk for development and progression of AD [9, 11, 28, 70]. Recent meta-analyses found that patients with insomnia had a 1.51-fold higher risk (95% CI: 1.06–2.14) of developing AD, [11] and that patients with sleep problems had a 1.55 (95% CI: 1.25–1.93) times higher risk of AD and a 1.65 (95% CI: 1.45–1.86) and 3.78 (95% CI: 2.27–6.30) times higher risk of cognitive impairment and pre-clinical AD, than people with no sleep problems, respectively [9].

However, evidence of the independent association of sleep problems and clinical outcomes is generally lacking in the literature. Caregiver burden is likely to be multifactorial and impacted by variables beyond patients' sleep symptoms, including the severity of patients' AD. Multiple patient and caregiver factors influence the institutionalization of AD patients, not only the overall clinical status of AD patients, but also the health, functionality, and economic status of their caregivers.

Nevertheless, insomnia and other sleep disturbances are recognized factors contributing to increased burden in AD patients, and the effective management and treatment of sleep symptoms may be helpful in improving clinical outcomes and delaying institutionalization. However, sleep problems are not only poorly characterized in this population, but there are also very few studies evaluating the effects of sleep treatments specifically in AD patients. Non-pharmacological sleep treatments, such as behavioral and light therapy, are typically used as first-line approaches in patients with insomnia and sleep disturbances, despite a lack of consistent data and standardized recommendations in this population [1, 75]. However, access to non-pharmacological sleep treatments is limited due to availability and

cost restraints [5]. The limited research for non-pharmacological sleep treatments reflects the obstacles faced with initiating practices in clinical care, particularly in low-resource environments, such as the cost, time taken to train carers, limited equipment, and lack of clear guidelines to deliver treatment [76]. For patients with dementia and in long term care, patients may also be unable or unwilling to partake in such therapies, preventing successful implementation [5]. Implementing approaches that actively include input from care staff, residents and their families may help overcome the challenges associated with non-pharmacological treatments in long-term care [76]. Practices that aim to improve the daily routines and quality of the sleeping environment for AD patients in care homes (such as the LOCK program, a specific frontline staff huddling program which enables staff to: **L**earn from bright spots; **O**bserve; **C**ollaborate in huddles; and **K**eep it bite-size) may also help alleviate sleep disturbances in this population [77]. Importantly, strategies that include interventions in several behavioral and environmental domains, such as sleep hygiene, light exposure and exercise, have shown some promise, which may suggest that a combined approach to behavioral therapy could improve outcomes [50]. Implementation of non-pharmacological treatments may reduce the likelihood of patient institutionalization and the associated economic burden, but successful implementation remains a key challenge in AD patients. Furthermore, many patients may require pharmacotherapies alongside these strategies to ensure patients can be treated at home for as long as possible.

Many currently available pharmacological treatments for insomnia are associated with a risk of further cognitive impairment, falls and fractures, and next-day residual effects in the elderly and therefore should be carefully evaluated prior to being prescribed. Studies investigating insomnia therapies in elderly patients have largely excluded patients with cognitive impairment or dementia. As a result, there remains uncertainty about the balance of benefits and risks associated with common sleep treatments when used in AD [78].

Few studies evaluating the use of sleep medications in AD patients were identified and were generally small. While treatments such as trazodone, [21, 40, 60] risperidone [53, 70] and suvorexant [35] showed some promise in improving clinical and caregiver symptoms, further data from larger, prospective trials are required. Previous reviews have drawn similar

conclusions, highlighting the scarcity and contradictory nature of current evidence [1, 79].

Benzodiazepines are also frequently used in AD patients, [79, 80] but a systematic literature review by DeFrancesco et al. (2015) reported no conclusive evidence for the improvements in sleep quality in AD patients taking benzodiazepines [81]. Moreover, benzodiazepines are associated with side effects including daytime sleepiness, worsened insomnia, confusion and amnesia. Though non-benzodiazepine ‘z-drugs’ may be associated with fewer side effects than benzodiazepines, [1, 10, 82] adverse events are still apparent. The American Geriatric Society recommends that both benzodiazepines and non-benzodiazepines be avoided in the elderly [14].

This review found mixed results between studies evaluating melatonin, which echoes findings from previous reviews [80, 83, 84]. A 2016 Cochrane review concluded that melatonin was unlikely to benefit AD patients with sleep problems [85]. Conversely, the combination of melatonin and bright-light therapy has been shown to have beneficial results in the nocturnal sleep of AD patients, [1, 10] suggesting a multifaceted approach targeting circadian rhythm stabilization may enhance the outcomes of sleep treatments in AD, which is not surprising given that AD patients tend to have both sleep and circadian rhythm disturbances.

The use of antipsychotics in the treatment of sleep and other behavioral disturbances in AD has been extensively discussed [1, 10, 82, 86, 87]. While they may be useful in the treatment of sleep disturbances and to alleviate neuropsychiatric symptoms, [70] they are not without risks. While no significant adverse events were reported in the studies included in this review, antipsychotics can cause residual sedation, confusion and falls, and have metabolic and cardiovascular side effects [1, 10, 86, 88, 89]. Antipsychotics also carry a black-box warning from the FDA for increased death in the elderly with dementia.

Suvorexant, currently the only approved pharmacotherapy for insomnia in AD, has demonstrated improved sleep and was well-tolerated in AD patients [35]. Suvorexant has also been shown to improve sleep maintenance and onset and was well-tolerated in elderly patients with insomnia [90]. Lemborexant, another orexin receptor antagonist, has also demonstrated promising results in improving sleep without significant adverse events in elderly patients [91]. Although not the focal population for this review, lemborexant has shown beneficial results in

nighttime and daytime symptoms with no worsening in cognitive functioning in AD patients with Irregular Sleep-Wake Rhythm Disorder (ISWRD) [92]. However, at present, the suvorexant trials in elderly patients and those with AD have reported far more conclusive results, based on objective PSG measures [35, 90]. While orexin receptor antagonists have been highlighted as promising options for elderly patients and those with dementia across the literature, data so far are limited [1, 79, 93].

The paucity of studies evaluating therapies in this population highlights the importance of further research in people with or at risk for AD, as there is insufficient evidence that treatments suitable for non-demented elderly persons will have an advantageous benefit-to-risk ratio in AD patients [14, 93]. Importantly many current studies do not adequately categorize sleep problems in AD patients which may include insomnia and circadian rhythm disorders. Successful treatment likely requires addressing both disorders which may have impacted the results in studies assessing the treatment of insomnia alone. As studies reporting data for sleep disorders other than insomnia, such as obstructive sleep apnea, were excluded from this analysis, this review has not discussed the burden of sleep disorders existing comorbidly with insomnia in AD patients. However, as the presence of comorbid sleep disorders may influence the clinical presentation and burden of symptoms, the impact of available sleep treatments in these patients should be further considered.

Limitations

There are several limitations to this review, with the most significant being that publications reported on sleep disturbances and insomnia without clear and consistent definitions. Sleep in AD is characterized by increased insomnia and sleep fragmentation, as well as circadian rhythm abnormalities, including deterioration and instability of rhythms, with delayed sleep phase and daytime napping. AD patients also have an increased rate of other sleep disorders that further contribute to sleep disturbance, including sleep apnea, sleep movement disorders (e.g., restless legs), and parasomnias such as REM sleep behavior disorder [94–97]. Furthermore, medications commonly given to AD patients including many antidepressants, cholinesterase inhibitors, and memantine can cause or worsen insomnia. In general, the studies covered in this review did not comprehensively assess and enumerate the specific sleep and rhythm disturbances

in participants, evaluate them for other sleep disorders, or control for medication effects. As a result, it is likely that sleep problems and sleep disorders other than insomnia were captured, particularly in studies that used non-specific definitions for sleep disturbance. For example, the NPI question used to assess sleep disturbance in many of the studies simply asks, “Does the patient awaken you during the night, rise too early in the morning, or take excessive naps during the day?”; it combines symptoms of insomnia with excessive sleepiness, and the questionnaire does not include screening for other sleep disorders.

Studies in this review generally did not control for other factors that could have influenced the clinical burden of disturbed sleep, most notably the degree of cognitive impairment and presence of comorbidities such as depression, agitation, and other behavioral disturbances commonly reported in AD patients. Thus, any associations, or lack thereof, may be obscured by the worse sleep and greater impairment that patients with severe AD inherently have.

Finally, only publications reporting specifically on AD patients were included and thus this review may have omitted data from studies on the burden of insomnia in AD patients which reported a population more broadly defined as dementia. Finally, the review and synthesis of the identified publications included only a qualitative review of the data and no further analyses were undertaken.

CONCLUSIONS

Insomnia/sleep disturbances confer a significant burden on both AD patients and caregivers and are an important risk factor for the institutionalization of patients. However, sleep and rhythm disturbances remain poorly characterized by clinicians and under-researched in this population, and no data were identified on the economic burden they may confer. Additional research is needed to fully quantify these clinical and economic burdens, especially given the large population at risk for AD. Furthermore, there is a paucity of data on effective treatments for insomnia in AD patients and the impact of treatment on clinical burden in patients and their caregivers.

The aging global population will increase the number of individuals at risk of developing AD. Sleep disturbances, particularly insomnia, appear to exacerbate this risk, and most currently available pharmacotherapies for insomnia have not been studied sufficiently in AD patients; treatment guidelines

advise against the use of many pharmacotherapies in older adults. Due to the impact of insomnia/sleep disturbances on AD risk and progression, and the associated patient and caregiver burden, there is a need to further evaluate the burden of disease in AD patients, investigate therapies targeted to this population, and determine whether treating insomnia and other sleep disorders may decrease the risk of developing AD and/or slow its progression.

ACKNOWLEDGMENTS

Medical writing assistance, under the direction of the authors, was provided by Ashley Enstone, Patrick Lavelle and Amy Sears of Adelphi Values PROVE™, in accordance with Good Publication Practice (GPP3) guidelines. This assistance was funded by Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc., Kenilworth, NJ, USA (MSD). Christopher Lines, PhD, and Alan Lipschitz, MD of MSD assisted with editing of the paper.

FUNDING

This study was supported by Merck Sharp & Dohme Corp., a subsidiary of Merck & Co., Inc., Kenilworth, NJ, USA.

SUPPLEMENTARY MATERIAL

The supplementary material is available in the electronic version of this article: <https://dx.doi.org/10.3233/JAD-215324>.

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