Sleep Disorders over the Full Range of Chronic Kidney Disease

Andreas Pierratos\textsuperscript{a}  Patrick J. Hanly\textsuperscript{b}

\textsuperscript{a}Department of Nephrology, Humber River Regional Hospital, University of Toronto, Toronto, Ont., and
\textsuperscript{b}Sleep Centre, Foothills Medical Centre, University of Calgary, Calgary, Alta., Canada

Introduction

Sleep disorders are common and under-recognized in patients at all stages of chronic kidney disease (CKD). A review of 17 studies in patients with end-stage renal disease (ESRD) indicated that ‘sleep disturbance’ was common with a mean prevalence of 44%. This review focuses on sleep apnea, insomnia, excessive sleepiness, restless legs syndrome (RLS) and periodic limb movement (PLM) disorder.

Sleep Apnea

Sleep apnea, in which patients stop breathing during sleep is classified as obstructive (OSA) due to intermittent closure of the upper airway or central due to intermittent loss of respiratory drive. Its severity is defined by the apnea-hypopnea index during a sleep study (polysomnography).

More than 50% of ESRD patients have sleep apnea \cite{1, 2}, which is dramatically higher than the general population (2–4%) and is similar in predialysis, peritoneal dialysis or hemodialysis patients \cite{1, 3}. The predominant type of sleep apnea found in patients with ESRD is OSA.

Sleep apnea is caused both by destabilization of central ventilatory control and upper airway occlusion during sleep. Enhanced ventilatory sensitivity to hypercapnia positively correlated with apnea severity \cite{4}, and a reduction in ventilatory sensitivity to hypercapnia following conversion from conventional hemodialysis (CHD) to nocturnal hemodialysis (NHD) was associated with a reduction in the severity of OSA \cite{5}. Upper airway occlu-
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sion can be caused by fluid overload and interstitial edema in the upper airway [6]. Displacement of fluid from the lower limbs increases neck circumference and pharyngeal resistance [7] and reduces upper airway cross-sectional area, contributing to the pathogenesis of OSA. Pharyngeal cross-sectional area measured with acoustic pharyngometry in patients on CHD was smaller than in controls, suggesting that this predisposes them to upper airway occlusion during sleep [8]. Conversion of ESRD patients from CHD to NHD was associated with an increase in pharyngeal cross-sectional area [9], possibly due to improved fluid removal. Conversion from continuous ambulatory peritoneal dialysis (CAPD) to nocturnal peritoneal dialysis has also been shown to reduce the frequency of OSA [10] associated with reduced pharyngeal narrowing as measured by magnetic resonance imaging [11]. Upper airway dilator muscle dysfunction due to neuropathy or myopathy associated with chronic uremia or the underlying cause of ESRD such as diabetes mellitus can also cause pharyngeal narrowing.

Advancing age, male sex and excessive weight are risk factors for the development of OSA in the general population. Some investigators have reported that these associations are weak in the ESRD population [1, 4, 12]. Obesity is not required for ESRD patients to develop OSA, and snoring is less intense than in patients with normal renal function.

Sleep apnea may exacerbate the symptoms of CKD such as daytime fatigue, sleepiness and impaired neurocognitive function. Sleep apnea may exacerbate the cardiovascular complications of ESRD. In patients with ESRD, hypoxemia during sleep is associated with nocturnal hypertension, left ventricular hypertrophy, impaired sympathovagal balance and an increased risk of cardiovascular complications. Coexisting sleep apnea may increase the risk of death in this patient population [13].

Although the characteristic clinical features of sleep apnea may be absent, a history of snoring, witnessed apneas during sleep and daytime sleepiness are suggestive of sleep apnea. Subjective sleepiness can be assessed with a number of simple scales, such as the Epworth Sleepiness Scale. Tools to estimate the likelihood of sleep apnea include the Berlin questionnaire and adjusted neck circumference, but these have not been validated in the ESRD population. Objective diagnostic testing can be limited to ambulatory monitoring at home (which typically records airflow, snoring, respiratory movement, oxygen saturation and heart rate). Alternatively, polysomnography can be performed in a sleep laboratory, which includes comprehensive monitoring of respiration, sleep and leg movements. Additional diagnostic tests such as the Multiple Sleep Latency Test (MSLT) and the Maintenance of Wakefulness Test can be considered for the evaluation of daytime sleepiness.

Sleep apnea should be treated if the patient has symptoms such as fragmented sleep and daytime sleepiness that are due to this disorder. In patients without sleep-related symptoms, consideration should be given to treating sleep apnea in those with severe disease (apnea-hypopnea index >30) since OSA of this severity has been associated with increased cardiovascular morbidity and mortality. Finally, sleep apnea should be treated if it is thought to be exacerbating co-existing medical disorders such as hypertension, myocardial ischemia and respiratory failure.

Treatment options include lifestyle modification (weight loss, avoidance of alcohol consumption close to bedtime, and avoidance of the supine position during sleep), the use of a dental appliance or continuous positive airway pressure to keep the upper airway open during sleep, and surgical correction of upper airway obstruction, such as removal of very enlarged tonsils, which is infrequent in adults. The experience from clinical practice suggests that ESRD patients have poor compliance with continuous positive airway pressure therapy, which is most likely due to the multifactorial etiology of sleep disruption in this patient population. Sleep apnea may be improved by changing the mode of renal replacement therapy. Although sleep apnea is not corrected by CHD or peritoneal dialysis, NHD, which enables patients to receive hemodialysis at home during sleep, has been shown to improve sleep apnea. Similar results have been reported in the CAPD population with nocturnal cycler-assisted peritoneal dialysis [10]. This was attributed to more effective fluid removal and its impact on the upper airway during sleep [11]. Although case reports have indicated correction of sleep apnea following successful kidney transplantation [14], results from case series are mixed, with some indicating that sleep apnea improves [15] and others indicating that sleep apnea resolves in only a minority of patients following transplantation [12].

Insomnia

Diagnosis is based on clinical assessment, with chronic complaints of difficulty initiating or maintaining sleep, waking up too early or unrefreshing sleep, despite adequate opportunity and circumstances for sleep, leading to impairment of daytime function.
Insomnia is common in patients with ESRD who are treated with CHD or CAPD (38–71%) [16–18] without a significant difference between the two.

The etiology of insomnia is multifactorial, and includes sleep disruption associated with RLS, PLM disorder and sleep apnea [19, 20], metabolic factors, bone pain and pruritus, psychiatric disorders such as depression, circadian rhythm disorders such as delayed sleep phase syndrome, medications, and poor sleep hygiene including frequent napping during daytime dialysis.

Polysomnographic studies have reported a total sleep time of 4.4–6 h per night, fragmented by a high frequency of arousals (up to 30/h), resulting in a sleep efficiency that ranged from 66 to 85% [19, 21, 22]. Stage 1 and 2 non-rapid eye movement sleep is often increased, whereas slow-wave sleep and REM sleep are reduced [19, 23].

Insomnia is a significant source of stress, which impairs quality of life [17] but has also been associated with shorter survival in patients on CHD and CAPD [24].

Clinical assessment and questionnaires can be used to quantify the severity of sleep-related symptoms. Objective sleep testing is generally reserved when comorbid sleep disorders such as sleep apnea, RLS and PLM, are suspected. The clinical assessment includes history of medical and psychiatric comorbidities, medications, alcohol and caffeine consumption. Completion of a 2-week sleep diary may help. Questionnaires such as the Pittsburgh Sleep Quality Index (PSQI), and the Insomnia Severity Index can be used. Objective testing of sleep can be performed by techniques including actigraphy, ambulatory monitoring and polysomnography.

Therapeutic options include: (1) correction of underlying medical conditions including ESRD, (2) nonpharmacologic strategies to consolidate nocturnal sleep, and (3) medications to improve sleep-related symptoms. There have been a few studies that have evaluated the impact of renal function replacement on insomnia. Novak et al. [25] found that sleep-related symptoms component of the Pittsburgh Symptom Score, improved significantly in 45 patients who were started on CAPD. An RCT comparing automated peritoneal dialysis with CAPD included a subjective description of ‘sleep quality’ [26], but no differences in measures of sleep quality were found. In a study which reported that NHD improved sleep apnea, there was no improvement in polysomnographic features of sleep fragmentation [19]. The prevalence of poor sleep, reflected by the PSQI, has been reported as 30% in a cohort of 115 patients with renal transplantation, but sleep quality before transplantation was not reported [27].

In a study of 24 CAPD patients, cognitive behavioral therapy was associated with a small but significant improvement in the PSQI score compared to the control group [28]. Although a variety of hypnotic and sedating medications are commonly used in the management of chronic insomnia, there have been no appropriately designed studies to evaluate their safety and efficacy in patients with ESRD.

Excessive Sleepiness

Excessive sleepiness is defined as the inability to stay awake or alert during the major waking episodes of the day, resulting in unintended lapses into drowsiness or sleep.

Sixty-six percent of patients in a CHD unit complained of daytime sleepiness [29]. Sleepiness appears to be equally prevalent in ESRD patients treated with CAPD as in those receiving CHD.

Hanly et al. [21] evaluated 24 unselected patients treated with CHD. Subjective sleepiness using the Epworth Sleepiness Scale (ESS <8) was reported in 50% of patients, and objective sleepiness using the MSLT (sleep latency <5 min) was found in 50% of patients. Sleepiness was associated with both a higher BUN and a higher frequency of PLM. Parker et al. [22] evaluated 46 selected CHD patients after excluding those with known causes of daytime sleepiness. Despite this screening, 50% of patients had sleep apnea and 50% had PLM. Thirteen percent of patients had evidence of ‘pathological’ sleepiness (sleep latency <5 min). Objective sleepiness correlated with indices of sleep apnea severity and the frequency of arousals on polysomnography [22]. The respiratory disturbance index, a measure of apnea frequency, accounted for only 11% of the variance in the measures of sleepiness, which implies that nonrespiratory factors play a significant role. In addition, 48% of patients had REM sleep on at least one nap, and 17% of patients had REM sleep on two or more naps, which may reflect the severity of night-time sleep disruption or may indicate a basic desynchronization of the timing of REM sleep.

Daytime sleepiness can be quantified clinically as falling asleep involuntarily in either passive (e.g. reading, watching TV) or active (e.g. driving, during conversation) situations. Sleepiness can be further assessed by questionnaires such as the Epworth Sleepiness Scale and the Stanford Sleepiness Score and by laboratory tests such as the MSLT and the Maintenance of Wakefulness Test.
The consequences of daytime sleepiness can range from decreased productivity at work to an increased risk of motor vehicle accidents.

Treatment of excessive sleepiness is primarily directed at the underlying cause, which can include clinical entities such as insufficient sleep, sleep disruption from a coexisting sleep disorder such as sleep apnea, RLS, PLM and narcolepsy, side effects of sedating medications, and chronic medical and psychiatric disorders. If the cause is unclear (idiopathic hypersomnolence) and the patient has disabling sleepiness, stimulant medications such as modafinil, methylphenidate and amphetamines may be considered. The impact of NHD on sleepiness in patients with ESRD has been evaluated [21]. Fifteen ESRD patients had overnight polysomnography and MSLT before and after they were converted from CHD to NHD. The data suggested that sleepiness improves following conversion to NHD, possibly related to reduced sleep disruption associated with PLM.

**RLS and PLM Disorder**

RLS refers to complaints of an unpleasant sensation in the legs accompanied by an urge to move them, typically in the evening or early part of the night, that is worse during periods of inactivity and transiently relieved by movement [30]. PLM disorder refers to involuntary jerking movement of the legs (and occasionally the arms) during sleep that may be disruptive to the patient or bed-partner. RLS is almost always associated with PLM disorder, but PLM can occur in the absence of RLS.

RLS has been reported in 14–23% in patients on CHD [31] and 20–57% in CKD patients [30]. The prevalence of PLM is greater than 50% in the CHD and CAPD population [19, 21, 22] associated with sleep onset and sleep maintenance insomnia [18].

The pathogenesis of RLS and PLM in patients with ESRD is not understood. Common risk factors include anemia, iron deficiency, elevated serum calcium and peripheral and central nervous system abnormalities. In addition, it is likely that alteration of dopamine and opioid activity in the nervous system plays a role. Correction of anemia by treatment with erythropoietin or intravenous iron was associated with a significant improvement in RLS and PLM. Peripheral neuropathy, secondary to uremia or the underlying cause of ESRD (such as diabetes) may also predispose patients to develop RLS and/or PLM.

RLS associated with ESRD appears to progress more rapidly, become more severe and is less responsive to dopaminergic medications than idiopathic RLS.

RLS is associated with difficulty initiating sleep, poor sleep quality and impaired health-related quality of life [31, 32]. Furthermore, RLS has been associated with depression in patients with CKD [33]. These sleep disorders have also been associated with increased mortality in patients with ESRD [32, 34].

RLS is diagnosed clinically. PLM disorder is diagnosed by polysomnography, which reveals periodic, involuntary movements of the legs during sleep. In order to determine whether PLM are responsible for complaints of sleep disruption, a trial of pharmacologic therapy may be required.

The indications to treat RLS and PLM are based on the severity of associated symptoms. There is no indication to treat asymptomatic PLM. General treatment measures include reduction of potential exacerbating factors such as excess caffeine, alcohol and nicotine, medical conditions (anemia, iron deficiency), and medications (tricyclic antidepressants, serotonin reuptake inhibitors, dopamine antagonists). Medications used to treat RLS and PLM include L-dopa and dopamine agonists such as pramipexole. These are favored over benzodiazepines such as clonazepam and opiates. Gabapentin may also be considered as an alternative medication. The frequency of PLM did not change when patients were converted from CHD to NHD [19]. However, kidney transplantation has been associated with an improvement in both RLS and PLM in several small studies [35, 36].

**Conclusion**

Sleep disorders are very common in patients with ESRD. Recognition and treatment of these disorders has the potential to both improve the quality of life and decrease morbidity and mortality in this patient population.

**Disclosure Statement**

The authors declare no conflict of interest.
References


