Obstructive sleep apnea: transition from pathophysiology to an integrative disease model

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Summary
Obstructive sleep apnea (OSA) is characterised by recurring episodes of upper airway obstruction during sleep and the fundamental abnormality reflects the inability of the upper airway dilating muscles to withstand the negative forces generated within the upper airway during inspiration. Factors that result in narrowing of the oropharynx such as abnormal craniofacial anatomy, soft tissue accumulation in the neck, and rostral fluid shift in the recumbent position increase the collapsing forces within the airway. The counteracting forces of upper airway dilating muscles, especially the genioglossus, are negatively influenced by sleep onset, inadequacy of the genioglossus responsiveness, ventilatory instability, especially post arousal, and loop gain. OSA is frequently associated with comorbidities that include metabolic, cardiovascular, renal, pulmonary, and neuropsychiatric, and there is growing evidence of bidirectional relationships between OSA and comorbidity, especially for heart failure, metabolic syndrome, and stroke. A detailed understanding of the complex pathophysiology of OSA encourages the development of therapies targeted at pathophysiological endotypes and facilitates a move towards precision medicine as a potential alternative to continuous positive airway pressure therapy in selected patients.

KEYWORDS
comorbidity, diagnosis, epidemiology, obstructive sleep apnea, pathophysiology, treatment

1 INTRODUCTION

Obstructive sleep apnea (OSA) is characterised by recurring episodes of upper airway obstruction during sleep, leading to markedly reduced (hypopnea) or absent (apnea) airflow at the nose and mouth. The condition is usually associated with loud snoring and intermittent hypoxaemia, and apneas are typically terminated by brief micro-arousals, which result in sleep fragmentation and diminished amounts of slow-wave sleep (SWS) and rapid-eye-movement (REM) sleep (Deegan & McNicholas, 1995). Patients with OSA are usually unaware of this sleep disturbance but the changes in sleep architecture contribute significantly to the prominent symptoms of unrefreshing sleep and excessive daytime sleepiness (EDS) typically reported by many of these patients (Levy et al., 2015). Furthermore, the intermittent hypoxaemia and sleep fragmentation associated with OSA generate cell and molecular responses that generate systemic inflammation, sympathetic excitation, and other responses that predispose to comorbidities, especially cardiometabolic and neuropsychiatric (McNicholas, 2019).

While anecdotal reports on breathing stoppages during sleep were already published long ago, a full account of the OSA syndrome was given for the first time by Guilleminault et al. in the 1970s. In several seminal papers, clinical manifestations and typical findings on
polysomnography (PSG) were described (Guilleminault et al., 1976). These authors and other independent research teams confirmed that previously unexplained medical conditions originate in respiratory problems occurring during nocturnal sleep. Current evidence indicates that OSA is common in the community and has a considerable impact on public health. Strong associations have been shown between OSA and drowsiness-related motor vehicle accidents (Bonsignore et al., 2019). There is accumulating evidence that untreated OSA is associated with hypertension and constitutes a risk for cardiovascular disease (McNicholas et al., 2007). For these reasons, adequate management of OSA proves a major concern for public health authorities.

2 | PATHOPHYSIOLOGY

While the detailed pathophysiology of OSA is complex, the fundamental abnormality reflects the inability of the upper airway dilating muscles to withstand the negative force generated within the upper airway during inspiration (Figure 1). In the normal setting, upper airway dilating muscles contract in a co-ordinated manner that is timed with each inspiration, thus counteracting the negative pressure that is generated within the upper airway during inspiration. Factors that increase this negative pressure or diminish the efficacy of dilating muscle contraction upset this balance and predispose to upper airway obstruction (Deegan & McNicholas, 1995). Narrowing of the upper airway increases upper airway negative pressure during inspiration, thus promoting collapse. Factors contributing to narrowing include craniofacial bony morphology, soft tissue accumulation from obesity or adenotonsillar hypertrophy, and transient factors such as fluid accumulation that gravitates towards the neck in the recumbent position.

2.1 | Upper airway narrowing

Most patients with OSA demonstrate a narrowed oropharyngeal airway that can be clinically assessed by the Mallampati score (Yu & Rosen, 2020), and genetic factors play a major role (Chi et al., 2014). Cephalometric and computed tomography (CT) studies demonstrate bony dimensions in the lower face and neck that result in upper airway narrowing (Neelapu et al., 2017), and clinical assessment demonstrates micro- or retrognathia in many of these patients (McNicholas, 2008a). Children with the Robin sequence or Treacher-Collins syndrome are especially prone to OSA because of bony changes to the lower face and/or mandible that result in structural narrowing of the oropharyngeal airway (Tan, Kheirandish-Gozal et al., 2016).

Soft tissue accumulation in and around the upper airway, such as with obesity and adenotonsillar hypertrophy, can predispose to OSA by narrowing the oropharyngeal lumen. Fat in the neck results in oropharyngeal narrowing and abdominal obesity reduces traction on the upper airway, which further predisposes to increased collapsibility (Deegan & McNicholas, 1995). Adenotonsillar hypertrophy is an important contributing factor in paediatric OSA, often in association with obesity (Dayyat et al., 2009).

Fluid accumulation in patients with congestive heart failure (CHF) and end-stage renal failure, predisposes to OSA by nocturnal redistribution of fluid in the recumbent position to the parapharyngeal soft tissues, which increases upper airway resistance and collapsibility (Lyons et al., 2017; White & Bradley, 2013).

Nasal obstruction, especially variable nasal obstruction as in rhinitis, contributes to the pathophysiology of OSA (McNicholas, 2008b; McNicholas et al., 1982) and intranasal corticosteroids have been reported to reduce the apnea–hypopnea index (AHI) in patients with rhinitis and mild to moderate OSA (Kielty et al., 2004). The supine posture also has an adverse effect on upper airway patency (Yildirim et al., 1991), largely due to gravitational forces.

2.2 | Upper airway dilator muscle function

Oropharyngeal airway patency is dependent on pharyngeal dilator muscles, especially the genioglossus, which act to stiffen the collapsible segment during inspiration (Deegan & McNicholas, 1995). These muscles contract in a phasic manner that is co-ordinated with inspiration, which precedes diaphragmatic contraction by milliseconds (Strohl et al., 1980). Activity of these upper airway muscles is modulated by chemical stimuli, vagal input, changes in upper airway pressure, and baroreceptor activity (Brouillette & Thach, 1980).

In OSA, a narrowed upper airway generates greater inspiratory collapsing force, which requires more forceful dilating muscle contraction to maintain airway patency. Dilating muscle activity is higher than normal subjects during wakefulness and diminishes to a greater extent during sleep, thus predisposing to obstruction (Mezzanotte et al., 1996), especially in REM sleep (Carberry et al., 2016). Overall, the deficit in OSA relates to inadequate compensation in the face of increased inspiratory negative pressure rather than a primary deficiency in muscle function, which is compounded by these being skeletal muscles, resulting in a greater decrement during sleep than the diaphragm.
2.3 | Respiratory control

The pattern of recurring apnea frequently observed in OSA supports an instability of ventilatory control similar to periodic breathing and upper airway obstruction is most likely when diaphragmatic and genioglossal inspiratory electromyogram (EMG) activity is at the lowest point of the cycle (Deegen & McNicholas, 1995). EMG activity progressively increases through the later stages of apnea and apnea termination is followed by hyperventilation for several breaths, after which both EMGs then decrease in activity, which predisposes to further obstruction (Dempsey et al., 2010).

2.4 | Apnea threshold

Normal subjects demonstrate fluctuations in ventilation associated with the transition from wakefulness to non-REM sleep, which is due to a reduction in the carbon dioxide (CO2) drive to breathe and the exposing of a sensitive apneic threshold that is critically CO2 dependent (Phillipson, 1978). In OSA, this threshold is amplified by post-apnea hyperventilation, resulting in CO2 reduction, and predisposing to further apnea (Dempsey et al., 2010). Additional factors that may contribute to further apnea post hyperventilation include lung stretch receptor and baroreceptor stimulation (Deegan & McNicholas, 1995).

2.5 | Loop gain

The predisposition to apnea associated with recurring cycles of hyper- and hypoventilation during sleep varies relating to loop gain, which refers to the gain of the negative feedback loop that regulates ventilation in response to a ventilatory disturbance. A high loop gain occurs where the magnitude of the increase in ventilation following apnea is high, thus increasing ventilatory system instability and increasing the likelihood of subsequent apnea (Dempsey et al., 2010). Two types of respiratory control system gain are evident, namely plant gain, which relates to the background drive to breathe, and controller gain, which relates to chemoreponsiveness (Dempsey et al., 2010). A reduced ventilatory drive and associated hypoventilation increases susceptibility to apnea by requiring only small transient ventilatory overshoots to reach the apneic threshold (high plant gain). Controller gain describes the slope of the ventilatory response to CO2 and an increased slope results in an increased susceptibility to apnea even in the setting of background hyperventilation and low plant gain.

2.6 | Arousal

Termination of apnea is usually associated with brain arousal, which may be an important protective mechanism (Eckert & Malhotra, 2008), but may also contribute to the pathophysiology of OSA by predisposing to further upper airway collapse because of increased post-apneic hyperventilation (Jordan et al., 2007; McNicholas, 1998). The intensity of respiratory cortical arousals appears to be a distinct pathophysiological feature that is associated with disease severity in patients with OSA (Bahr et al., 2021). Increasing ventilatory effort appears to be the most important arousal stimulus (Deegan & McNicholas, 1995).

The arousal response varies in patients with OSA and can be quantified by the arousal threshold, which can be assessed noninvasively by PSG (Sands et al., 2018b). A low arousal threshold is an important potential contributing factor to OSA pathophysiology and may represent a therapeutic target in selected patients (Eckert et al., 2011).

2.7 | Integrated pathophysiology and implications for treatment

The relevance of physiological, non-anatomical factors in the pathophysiology of OSA has generated major interest in recent years (Randerath et al., 2018) and in one study of subjects with and without OSA, similar proportions of subjects, roughly one-third each, had the endotypic traits of a minimal genioglossus muscle responsiveness during sleep, a low arousal threshold, or a high loop gain, with 28% of subjects having more than one of these traits (Eckert et al., 2013).

While the basic deficit of increased upper airway collapsibility in OSA can be readily reversed by continuous positive airway pressure (CPAP) therapy, a detailed understanding of the pathophysiology opens the potential for other management options (Schutz et al., 2021). Inadequate upper airway dilating muscle compensation may be improved by targeted pharmacotherapy (Taranto-Montemurro et al., 2019). Sleep-induced reduction in respiratory motor neurone output can be reversed by electrical stimulation of the hypogossal nerve (Strollo et al., 2014). Acetazolamide may benefit OSA in selected patients with a high loop gain (Edwards et al., 2012) and zolpidem increases sleep efficiency and the respiratory arousal threshold (Messineo et al., 2020).

3 | Diagnosis

The high global prevalence of OSA (Benjafield et al., 2019) represents a challenge for diagnosis as the current “gold standard” diagnostic test is PSG in a sleep laboratory, which is labour intensive and expensive (Kapur et al., 2017). Disease severity is currently measured by the AHI as determined from a sleep study. However, there is a poor relationship between daytime symptoms such as sleepiness and the severity of OSA recorded in a sleep study (Deegan & McNicholas, 1996), and there is a growing trend to move away from the AHI as the principle measure of OSA severity towards a more personalised approach to OSA diagnosis and treatment (Pevernagie et al., 2020), by considering individual risk factors, clinical history and comorbid disease in the diagnosis and treatment of OSA (Randerath et al., 2018). Other signals such as heart rate variability, pulse transit time, expanded analysis of the oximetry signal, and the use of biomotion sensors may
allow for an improved and phenotypic diagnosis (McNicholas, 2021). Home sleep apnea testing (HSAT) and wearable technologies may help improve access to diagnosis (O’Mahony et al., 2020), and advances in telemedicine help to strengthen inter-departmental collaboration, thus improving the overall care of patients with OSA (O’Donnell et al., 2020).

Patients with OSA typically present with EDS, frequent awakenings, bed partner reports of frequent choking/gasping during sleep, and loud snoring (Kapur et al., 2017), in addition to morning headache, dry mouth, nonrestorative sleep, and may have hypertension or other comorbidity (McNicholas, 2008a). However, many patients do not report EDS or fatigue with some patients reporting only minimal symptoms (Heinzer et al., 2015), which have important implications for management and disease outcomes (Gagnadoux et al., 2016).

The Epworth Sleepiness Scale (ESS) is the most widely used clinical tool to evaluate subjective sleepiness, but correlates poorly with AH1 (Kingshott et al., 1995) and is open to reporting bias. Daytime symptoms in OSA are influenced by age, gender, and the presence of other comorbidities (Levy et al., 2015), particularly depression and insomnia, and other symptoms such as fatigue and tiredness may be equally important in certain groups. Anthropometric and other objective variables such as age, sex, body mass index (BMI), neck circumference and co-morbidities may be more reliable than subjective variables such as snoring and EDS in predicting OSA (Ustun et al., 2016). The identification of clinically significant OSA may be improved by the inclusion of relevant comorbidities, especially the loss of nocturnal blood pressure (BP) dipping (Crinion et al., 2017).

3.2 | Home sleep apnea testing

Home sleep apnea testing can be performed in the patient’s home, is cheaper and more convenient than PSG. HSAT will record between four and seven variables that include respiratory effort, airflow, heart rate or electrocardiograph (ECG), arterial oxygen saturation, snoring, body position and movement. Newer devices have improved diagnostic accuracy and have been validated against PSG (Collop et al., 2011). Also, HSAT can record several nights of sleep, which is advantageous as night-to-night variability is a feature of OSA and is most relevant in patients with mild disease (Guerrero et al., 2014). However, HSAT devices underestimate the AH1 due to their inability to record total sleep time and detect arousals (Escourrou et al., 2015). Fewer physiological variables are measured which can miss co-existing sleep disorders such as insomnia, periodic limb movements, and parasomnias.

3.3 | Clinical and pathophysiological phenotypes

Clusters of different clinical phenotypes can be identified among the broad population of patients presenting for assessment of OSA (Bailly et al., 2020). Furthermore, certain pathophysiological traits that are very common in OSA such as loss of nocturnal dipping of BP have significant implications for the development of associated comorbidity (Crinion et al., 2017). Thus, whatever sleep study is employed in the assessment of suspected OSA, the findings must be integrated into the overall assessment of the patient as regards clinical significance, and management should be linked to the underlying phenotype where symptom profile and additional factors to the AH1 such as acute systemic effects and associated relevant comorbidity are factored into the decision-making process (Randerath et al., 2018).

3.4 | New technologies

New measurements and technologies may better assess the various pathophysiological mechanisms underlying OSA, such as loop gain, arousal threshold, and anatomical factors. Reliable techniques to monitor sleep structure and arousals outside conventional EEG, such as arterial tonometry, require further evaluation and development, in addition to assessment of autonomic state and cardiovascular events associated with OSA events, such as ECG algorithms, heart rate variability, arterial tonometry, and capnography, which are currently omitted from conventional PSG (Khoo & Chalacheva, 2016). Further development of existing signals, such as cordless portable acoustic devices, allow enhanced diagnostic potential (Abbasi, 2017; Alshaer et al., 2016). There is a need to identify and validate physiological signals during wakefulness or sleep that may be useful in predicting cardiovascular risk.

The potential role of new technologies, like smartphone-based applications (Ko et al., 2015) and consumer-focused wearable devices (Haghayegh et al., 2019), are not yet sufficiently elaborated, and healthcare systems often do not integrate objective information...
Three-dimensional model of obstructive sleep apnea (OSA) disease severity. The x-axis represents the amount of respiratory events (A) in an overnight sleep period. While this number is usually expressed as the apnea–hypopnea index in clinical studies, the absolute count of events is considered in this model. The y-axis represents an acute systemic effect (E) induced by a respiratory event, e.g. a certain degree of hypoxaemia. The hatched area A*E represents the integration of all events with their associated systemic effects, being the nightly exposure to adverse effects of OSA. The nightly exposure inflicts repetitive strain to the end-organ systems. The z-axis represents a chronic end-organ impact (O) of OSA, e.g. cognitive impairment, arterial hypertension, cardiovascular damage, insulin resistance, etc. Due to interindividual differences in susceptibility, the end-organ impact may be variable among patients with OSA for similar levels of exposure. The dotted volume A*E*O represents the relationship between the three dimensions. Based on postulated differences in susceptibility, the disease spectrum may vary from low exposure/high impact to high exposure/low impact. The dashed aspect of the boundaries indicates that the end-organ impact of OSA is as yet difficult to assess. This is due to uncertainty regarding susceptibility on the one hand and to possible confounding effects of other disease processes on the other. Figure reproduced from (Randerath et al., 2018) with permission from the publisher.

4 | COMORBIDITIES

Obstructive sleep apnea is independently associated with many comorbidities, including cardiovascular, metabolic, renal, and neuropsychiatric (McNicholas, 2019). However, many studies were cross-sectional in design, which limits the ability to evaluate causality, and there is growing evidence of a bi-directional relationship between OSA and comorbidity (Gleeson & McNicholas, 2022). Furthermore, most reports evaluating links between OSA and comorbidity have used the AHI as the principal metric of OSA severity, and there is growing evidence that the relationship of OSA with comorbidity should take into account additional variables such as acute and chronic systemic effects (Randerath et al., 2018). Figure 2 illustrates the theoretical relationship between respiratory events, their systemic effects, and potential end-organ impact.

4.1 | Cardiovascular disease

Obstructive sleep apnea is proposed as an independent risk factor for cardiovascular disease (McNicholas et al., 2007) and potential mechanisms include intermittent hypoxaemia (Ryan et al., 2005), sleep fragmentation (Carreras et al., 2014), exaggerated intrathoracic pressure swings (Bradley et al., 2001), sympathetic excitation (Jullian-Desayes et al., 2015), inflammation (Ryan et al., 2009), and oxidative stress (Eltzschig & Eckle, 2011). However, these potential mechanistic relationships are complex, and, for example, there is evidence that mild hypoxaemia may be cardioprotective (Almendros et al., 2014). Furthermore, randomised controlled trials (RCTs) of positive pressure therapy of OSA have failed to show convincing evidence of benefit in the secondary prevention of cardiovascular disease (McEvoy et al., 2016; Peker et al., 2016), which challenges the notion that OSA is an independent contributor to cardiovascular morbidity. As further explained in the treatment section below, the relationship of OSA to cardiovascular disease and its outcomes remains uncertain.

4.1.1 | Hypertension

Obstructive sleep apnea is a risk factor for systemic hypertension, often with a non-dipping nocturnal BP profile (Parati et al., 2013), including data from the Sleep Heart Health Study (Nieto et al., 2000) and the Wisconsin Sleep Cohort Study (Peppard et al., 2000). A meta-analysis of seven prospective studies confirmed an independent relationship with incident hypertension in a dose-dependent fashion (Xia et al., 2018), which is often associated with resistance to conventional anti-hypertensive treatment (Hou et al., 2018). Indeed, non-dipping nocturnal BP is highly predictive of OSA, independent of symptom profile (Crinion et al., 2019), and REM-associated OSA is independently associated with incident non-dipping BP (Mokhlesi et al., 2015). Sympathetic excitation appears to be the principal pathogenic mechanism with possible additional involvement of renin–angiotensin–aldosterone...
system dysfunction (Crinion et al., 2021). Incident hypertension has also been independently associated with mild OSA, especially in patients aged <60 years (Vgontzas et al., 2019).

Continuous PAP therapy reduces BP, especially in younger subjects and those with uncontrolled hypertension or severe oxygen desaturation (Pengo et al., 2020), and in more CPAP-compliant patients (Levy & McNicholas, 2013). CPAP may also restore the nocturnal dipping pattern in normotensive patients with OSA (Crinion et al., 2021; Sapina-Beltran et al., 2019).

4.1.2 | Congestive heart failure

Congestive heart failure has a bi-directional relationship with sleep disordered breathing and is associated with both OSA and central sleep apnea (CSA). Typically, CSA in CHF is characterised by a central breathing pause that occurs during the decrescendo portion of the cyclic respiratory pattern (Naughton et al., 1993). CSA is associated with increased risk of incident heart failure (Gottlieb et al., 2010), which is proportional to OSA severity, and may be linked to major swings in intrathoracic pressure resulting in increased cardiac preload and afterload (Nicholl et al., 2014). The severity of intermittent hypoxaemia is a stronger predictor of outcome in patients with CHF than the AHI (Watanabe et al., 2017). CHF predisposes to OSA by nocturnal redistribution of fluid in the recumbent position to the parapharyngeal soft tissue, which increases upper airway resistance and collapsibility (White & Bradley, 2013).

4.1.3 | Coronary artery disease

The evidence for OSA as an independent risk factor for coronary artery disease is equivocal and is stronger for males than females (Dong et al., 2013; Gottlieb et al., 2010; Hla et al., 2015). Furthermore, CPAP therapy has not been associated with improved outcomes in patients presenting with acute coronary syndrome (Sanchez-de-la-Torre et al., 2020), except in those presenting with a first episode (Zapater et al., 2020).

4.1.4 | Atrial fibrillation (AF)

Obstructive sleep apnea is a significant risk factor for the development and recurrence of AF (Linz et al., 2018) and international guidelines for the management of AF recommend diagnosis and treatment of OSA (Calkins et al., 2018; Kirchhof et al., 2016), as untreated disease has been shown to reduce the efficacy of both pharmacological and catheter-based anti-arrhythmic therapy. A meta-analysis of seven prospective cohort studies found that CPAP therapy was associated with a reduction in AF recurrence, irrespective of whether they underwent pulmonary vein isolation (Shukla et al., 2015).

4.1.5 | Stroke

The relationship between OSA and stroke is complex and bi-directional (Bassetti et al., 2020). After adjusting for potential confounders such as age, sex, BMI, smoking, hypertension and diabetes, untreated OSA conveys a two-fold increased risk of stroke incidence (Yaggi et al., 2005), and this increased risk is confirmed by large prospective population studies such as the Sleep Heart Health Study (Redline et al., 2010) and the Wisconsin Cohort Study, which also reported that 25% of strokes occurred during the night (Young, 2009). The SAVE trial (ClinicalTrials.gov Identifier: NCT00738179) reported no reduction in stroke rate with CPAP therapy (McEvoy et al., 2016), but a more recent post hoc analysis of this trial identified self-reported snoring as a risk factor for incident stroke (Li et al., 2020), and CPAP compliance of >4 h may provide benefit (Parra et al., 2015). Randomised trials support an improvement in both short- and long-term functional outcomes with CPAP therapy in patients with OSA after stroke (Brill et al., 2018). Stroke may predispose to OSA by impairment of breathing control mechanisms at a central level and/or adversely affecting upper airway muscle function, and OSA is highly prevalent after stroke (Alexiev et al., 2018).

4.2 | Diabetes mellitus

Obstructive sleep apnea is independently associated with diabetes mellitus (Reutrakul & Mokhlesi, 2017). Several cross-sectional cohort studies, including the European Sleep Apnea Database (ESADA) (Kent et al., 2014a, 2014b), have demonstrated an independent relationship with type 2 diabetes and insulin resistance, which are also supported by long-term follow-up studies (Kendzerska et al., 2014; Lindberg et al., 2012). Potential mechanisms of diabetes and insulin resistance include intermittent hypoxaemia and sleep fragmentation leading to sympathetic excitation and inflammation (Reutrakul & Mokhlesi, 2017).

Some consequences of diabetes mellitus could predispose to OSA, including neuropathy affecting the upper airway muscles, and disturbances in ventilatory control. However, the impact of CPAP therapy on glycaemic control is uncertain (Chirinos et al., 2014; Reutrakul & Mokhlesi, 2017).

4.3 | Renal dysfunction

Renal disease and OSA have a bi-directional relationship (Gleeson & McNicholas, 2022). OSA is up to 10-times more prevalent in patients with chronic kidney disease (CKD) than the general population (Hanly, 2004), and potential contributing factors include increased chemoreflex sensitivity, reduced clearance of uraemic toxins, and hypervolaemia (Abuyassin et al., 2015). Fluid accumulation with associated nocturnal redistribution in the recumbent position, similar to CHF, is a key factor in end-stage renal disease, as evidenced by the beneficial impact on OSA of fluid removal during dialysis (Lyons et al., 2015).
Obstructive sleep apnea can also contribute to CKD (Ahmed et al., 2011), and possible mechanisms include hypertension and intrarenal hypoxaemia with glomerular hyperfiltration (Abuyassin et al., 2015). However, retrospective data analysis from the Wisconsin Sleep Cohort reported that OSA did not accelerate kidney function decline over time (Canales et al., 2018) and long-term follow-up of CPAP therapy in patients with OSA have shown no sustained benefit to renal function (Loffler et al., 2017; Rimke et al., 2021).

### 4.4 Depression

Depression and OSA may exhibit similar symptoms, including poor concentration, memory, and fatigue, which complicate their clinical assessment and diagnosis. The prevalence of depression in OSA ranges from 20% to 40% (Bixler et al., 2017) and there appears to be an increased odds ratio of depression with increasing severity of SDB (Peppard et al., 2006). A meta-analysis of 22 RCTs reported that therapy of OSA resulted in a dose-dependent improvement in depressive symptomatology using recognised depression scales (Povitz et al., 2014).

### 5 Epidemiology

The first major epidemiological study on OSA was published in 1993. Using PSG, Young et al. assessed the prevalence of OSA in the Wisconsin Sleep Cohort, a representative sample from the employed population (Young et al., 1993). The estimated prevalence of OSA, defined as an AHI of ≥5 events/h, was found to be 24% in men and 9% in women. Based on the combined presence of EDS and an increased AHI, it was estimated that 2% of women and 4% of men met the minimal diagnostic criteria for OSA disorder. The relevance of having OSA without EDS, which occurred in the majority of subjects, was not further addressed. Also, it was not explored whether OSA and EDS were causally related. To test this relationship, an interventional study would have been required.

Participants from the abovementioned cohort were invited for repeat studies at 4-year intervals. In a subsequent investigation, the data from the 1988–1994 period were compared with the 2007–2010 period, and the prevalence of OSA was assessed based on an AHI of ≥15 events/h, representing moderate-to-severe OSA (Peppard et al., 2013). The prevalence rates of moderate-to-severe OSA in the latter period were 10% among men aged 30–49 years; 17% among men aged 50–70 years; 3% among women aged 30–49 years; and 9% among women aged 50–70 years. As in the foregoing study, the prevalence of an increased AHI was higher than the prevalence of EDS. A direct comparison between both studies was not feasible because of differences in assessment of subjective sleepiness. Substantial increases in OSA prevalence were seen over the span of two decades, ranging between 14% and 55% depending on the subgroup. The rise in OSA figures was ascribed to the surge of prevalent obesity.

The HypnoLaus study is a large ongoing survey on the prevalence of OSA in the region of Lausanne, Switzerland (Heinzer et al., 2015). As in the latest Wisconsin study, an AHI of ≥15 events/h is used for identifying OSA. However, the method used for hypopnea scoring is different in the study design of these cohorts, the Wisconsin study applying a 4% desaturation criterion and the HypnoLaus study being compliant with the 2012 American Academy of Sleep Medicine (AASM) scoring criteria (Berry et al., 2012). Heinzer et al. found that the prevalence of OSA in their population-based sample was 23.4% in women and 49.7% in men. It was suggested that this high rate might be attributable to the increased sensitivity of the applied recording techniques and scoring criteria. Indeed, the 2012 AASM scoring system is much more prone to detecting hypopneas and thus produces higher AHI values. In keeping with the American studies, only a minority of men and women with an AHI of ≥15 events/h had concomitant EDS. In a subsequent analysis, Heinzer et al. applied prevailing AASM criteria of an AHI of ≥5 events/h plus symptoms and/or comorbidities (AASM, 2014; Heinzer et al., 2016) to test the occurrence of OSA in their population (Heinzer, Marti-Soler, & Haba-Rubio, 2016). The estimated prevalence was 79.2% in men and 54.3% in women. It was concluded that these overwhelming figures were probably not indicative of an OSA epidemic, but rather pointed to the unrealistic definition of OSA issued by the AASM.

Recently, a literature search was performed to try and estimate the global prevalence of OSA in individuals aged 30–69 years (Benjafield et al., 2019). The AASM 2012 scoring criteria were used as a reference for assessing the AHI, and a conversion algorithm was created for studies that had applied other criteria to allow determination of equivalent AHI values. Reliable prevalence data from 16 countries were extrapolated to other countries. Adjustments were based on population similarities, considering distributions of BMI, race, and geographical proximity. It was estimated that, globally, 936 million men and women of the considered age group have at least mild OSA and 425 million have moderate-to-severe OSA. The prevalence was highest in China, followed by the USA, Brazil, and India.

The use of the AHI as a primary predictor for identifying clinically relevant OSA in the general population is questionable. In all demographic studies to date, the prevalence of an increased AHI proves significantly higher than the associated prevalence of self-reported EDS. Whether people from the community in whom an increased AHI is found are at risk of having or developing OSA-related medical problems has not been established. If this risk would be substantial, the implementation of OSA screening programmes would be justified. The US Preventive Services Task Force have investigated this issue and have advised against population surveys for OSA in adults who are asymptomatic or who have unrecognised symptoms (Bibbins-Domingo et al., 2017). The task force was unable to determine the magnitude of the benefits versus harms of screening for OSA. For case finding to be appropriate, the essence of OSA as a clinically relevant disease must be defined beyond the AHI. With the introduction of a valid clinical definition, biases in future epidemiological studies could be avoided (Holley & Phillips, 2018).
6 | PHENOTYPES

Endotypes and phenotypes of SDB have been systematically studied in recent years (Edwards, Redline, Sands, & Owens, 2019). An “endotype” represents a particular mechanism that causes a physiological or metabolic disturbance in certain organ systems. As explained in the pathophysiology section above, several endotypes, in the sense of pathophysiological traits, have been distinguished in OSA. Evidence is accumulating that physiologically targeted treatment for OSA may effectively decrease the AHI and thus lower the pathophysiological burden of the disorder (Eckert, 2018).

While the term “phenotype” can have different meanings (and is sometimes used to also denote “endotype”), it usually refers to a combination of disease characteristics, in relation to clinically meaningful attributes (symptoms, treatment response, health outcomes, quality of life) that can be used to distinguish certain categories of patients from others (Zinchuk et al., 2017). Because OSA embodies a complex and heterogeneous disorder, several cluster analyses have been performed to discern clinical subtypes (Zinchuk & Yaggi, 2020). While certain profiles emerge from these studies, the method for subtyping clinical features is as yet not suitable to predict clinical outcomes in individual patients.

Originally, three symptomatic forms of OSA have been observed, namely patients with disturbed sleep, minimal symptoms, and EDS (Ye et al., 2014). These findings have been reproduced and expanded in a subsequent international, multicentric study (Keenan et al., 2018). In addition to the three basic clusters two other categories were recognised, i.e., upper airway symptoms dominant and sleepiness dominant subtypes. Similar average AHI values were found in both study samples and across clusters indicating that clinical phenotypes cannot be differentiated by the AHI. While the AHI fell short in predicting cardiovascular outcomes, a multisite study in a US Veteran cohort demonstrated that certain physiological endotypes, assessed by PSG, captured risk of adverse cardiovascular outcomes (Zinchuk & Yaggi, 2019).

Regarding cardiovascular outcomes, mixed results have been shown. There is evidence that the sleepy OSA phenotype is associated with worse clinical outcome and higher comorbidity (Randerath et al., 2018) and other phenotypes such as non-dipping nocturnal BP have a high diagnostic prediction and are associated with a higher risk of comorbidity (Crimion et al., 2019). A recent survey from the USA seems to confirm the cardiovascular implications of EDS in OSA. Using data from the Sleep Heart Health Study, the excessively sleepy subtype was found to be most strongly associated with prevalent heart failure and with incident cardiovascular disease (Mazzotti et al., 2019). In contrast, the disturbed sleep or “insomnia” subtype, a predominant cluster in a survey of ESADA, was more frequently linked with cardiovascular comorbidity than the sleepy phenotype, despite lower mean AHI values (Saaresranta et al., 2016).

Of note, OSA is not only a complex and heterogeneous disorder, but may also co-occur with other highly prevalent sleep disorders (Pepin et al., 2018). Unhealthy lifestyle habits and manifestations of medical and psychiatric comorbidities may further blend in with the palette of symptoms, thereby generating mixed phenotypes. The challenge for future research lies in the identification of unique markers of OSA to estimate its relative contribution in these composite conditions, and to predict response to targeted treatment more reliably (Edwards et al., 2019; Pevernagie, 2021).

7 | TREATMENT

The aim of treating SDB is to maintain the patency of the upper airway, to stabilise the breathing pattern and to ensure adequate ventilation of the lungs. Because OSA and CSA frequently co-occur, and share clinical and pathogenetic features (Dempsey et al., 2014), certain therapeutic strategies are common to both.

Anatomical traits compromising the patency of the upper airway are predominant in OSA. These traits are related to the surrounding musculoskeletal structure, body mass and muscular responsiveness. Non-anatomical traits including arousability and loop gain may be present in both OSA and CSA. Habits such as alcohol use, sleep medication, lifestyle, body position while sleeping may influence both anatomical and non-anatomical traits and thus the severity of the breathing disturbance. Part of the treatment plan is to address these variables, by which the AHI can readily be improved (Kuna et al., 2013; Strobel & Rosen, 1996).

Supplemental oxygen has been used for treating respiratory sleep disorders with varying degrees of success. Nocturnal oxygen therapy (NOT) may be beneficial for treating CSA with Cheyne-Stokes respiration in heart failure (Aurora et al., 2012). Increasing the arterial PO2 is thought to dampen the respiratory drive, thus reducing minute ventilation and increasing CO2 reserves (Bordier et al., 2016). While supplemental oxygen is not recommended as a first-line treatment for OSA, it may be co-administered to reduce residual hypoxaemia in patients using CPAP. Promise lies in the identification of NOT-responsiveness in patients with OSA, based on the assessment of endotypic traits using diagnostic PSG (Sands et al., 2018a). Combination of NOT with non-CPAP treatment modalities might also improve clinical outcomes (Edwards et al., 2016).

Several pharmacological agents have been tested in different types of SDB. While some drugs may target treatable traits of OSA or CSA, there are as yet no compounds that are licensed in this domain (Gaisl et al., 2019). Acetazolamide, a carbonic anhydrase inhibitor, is known to reduce loop gain of the ventilatory control (Edwards et al., 2012), making it suitable for treating both CSA and OSA. A meta-analysis of nine studies in patients with heart failure and CSA confirmed a modest, but statistically significant decrease of AHI (Wongboonsin et al., 2019). In OSA, the AHI may also be lowered, but daytime sleepiness and other clinical outcomes have not been shown to improve (Gaisl et al., 2019). Because long-term studies with acetazolamide are lacking, its role in the extended treatment of respiratory sleep disorders is uncertain.

For a long time, hypnotics were believed to have adverse effects on respiration during sleep. Recent studies shed a different light on this conception. The effects of hypnotics in CSA are not well-known,
although some beneficial effect on the AHI may be anticipated due to inhibition of arousal and increase of CO₂ stores. In OSA, interest was revived in the therapeutic potential of drugs that suppress arousability and that might reduce the AHI in patients with OSA with a low arousal threshold (Eckert et al., 2011). Against expectation, zolpidem was not found to increase the AHI, but to promote a positive effect by stimulating genioglossus activity during sleep in patients with OSA (Carberry et al., 2017). Due to a lack of randomised trials, no recommendations can presently be made regarding the selection of patients with OSA who could benefit from treatment with hypnotic drugs (Carberry et al., 2018).

The use of devices that deliver PAP to the airways is the mainstay of treating OSA (Sullivan et al., 1981). Besides mechanically stabilising the upper airway, PAP devices may have other effects on the respiratory system, such as increasing the end-expiratory lung volume and improving stability of the central respiratory drive. These mechanisms may also be beneficial in heart failure with CSA and Cheyne-Stokes respiration, as some of these patients are responsive to CPAP treatment (Bradley et al., 1999). Problems with acceptance and tolerance may compromise CPAP treatment compliance. Adequate compliance, defined as CPAP use for ≥ 4 h/night for ≥ 70% of nights, was reported insufficient in >40% of patients with OSA in the initial treatment phase (Weaver & Grunstein, 2008). Many factors determine compliance with CPAP therapy. OSA disease severity and symptomatic benefit predict better outcomes, whereas adverse physical and psychological effects are linked with unsatisfactory results. Patient education and supportive interventions are paramount to treatment success (Smith et al., 2009). However, absence of any symptomatic improvement despite sufficient compliance should prompt considering other diagnoses and may justify discontinuation of CPAP therapy.

Mandibular repositioning devices (MAD) are increasingly used for treating primary snoring and OSA (Ramar et al., 2015). This therapy has been proven effective in patients with OSA who have a mildly to moderately elevated AHI. Healthy dentition is required, as the appliance must be anchored to the teeth of the upper and lower jaw. MAD therapy generally reduces the AHI and improves EDS. In studies comparing CPAP and MAD therapy, the effects on the AHI and EDS were better in the former group, but seemed similar in patients with milder OSA receiving either CPAP or MAD (Sharples et al., 2016).

Last but not least, surgical procedures for OSA can be applied in carefully selected patients. For a thorough review of these procedures the reader is referred elsewhere (Pevernagie et al., 2021). Below is a concise description of the most common interventions.

- Nasal surgery aims at improving the endonasal lumen to facilitate nasal breathing. It usually has only a minor effect on OSA severity, but may improve compliance to PAP therapy in patients with OSA with nasal obstruction (Randerath et al., 2011).
- Pharyngeal surgery comprises resection of adenoids and/or tonsils when these structures are enlarged, as well as remodelling of pharyngeal structures in patients with OSA, preferably selected after drug-induced sedation endoscopy (Rashwan et al., 2018).
- Hypoglossal nerve stimulation is a relatively new surgical technique based on electrical stimulation of a branch of the hypoglossal nerve that induces tongue protrusion. A set of components is implanted, including a pulse generator, a sensor lead monitoring inspiration and a stimulation electrode leading to the hypoglossal nerve. Candidates for this treatment option are symptomatic patients with OSA who are intolerant to PAP therapy (Strollo et al., 2014).
- In patients with cranio-skeletal deficiencies, maxillomandibular advancement surgery may offer a permanent remedy for OSA. The procedure consists of a bilateral sagittal split osteotomy of the mandibula and a Le Fort I osteotomy of the maxilla. Both structures are advanced by ~10 mm and fixed in this new position. Patients with OSA who have been appropriately selected for this
intervention may have a lasting improvement in EDS and AHI, among other variables (Camacho et al., 2019).

Finally, bariatric surgery can be an appropriate remedy for obese patients with OSA in whom dietary regimens have failed. Overall, bariatric surgery is successful in reducing weight and the AHI, but in some patients persistent symptoms of OSA may prompt continuation of specific treatment for OSA (Greenburg et al., 2009).

**8 | FUTURE CHALLENGES**

Despite extensive knowledge on the pathophysiological mechanisms and the adverse systemic effects of SDB, only small and/or nonsignificant results from CPAP therapy have been demonstrated in clinical trials aiming to reduce the cardiovascular burden in patients with OSA. These findings are paradoxical and prompt further consideration. It can no longer be denied that problems with the basic concept of OSA must be addressed as an obligatory step for respiratory sleep medicine to advance. Two aspects should be considered in particular: the causality issue and differences in individual susceptibility of systemic effects of OSA.

It is commonly assumed that the co-occurrence of symptoms and signs suggestive of OSA with an increased AHI implies a causal relationship. However, many symptoms and signs of OSA are nonspecific (e.g., disturbed sleep, EDS, fatigue, high BP) and may fit other diagnoses as well (e.g., insomnia, narcolepsy, sleep restriction, primary hypertension). Therefore, the association between clinical manifestations and pathophysiological findings in OSA may due to coincidence (Figure 3). In that instance, treatment of OSA will be ineffective. Symptomatic response to therapy is a factor that corroborates causality, but treatment responsiveness is as yet difficult to predict.

Another plausible explanation for equivocal therapeutic results may lie in differences in individual susceptibility to systemic effects of OSA (Pevernagie et al., 2020; Randerath et al., 2018) (Figure 2). Some subjects may be vulnerable, whereas others may be relatively resistant, irrespective of the AHI value or even the degree of hypoxaemia.
Thus, disease severity may be disparate among subjects with similar levels of exposure. Consequently, results of treatment may average out to naught as categories with various degrees of disease severity (despite equivalent AHI or ODI values) are being recruited in clinical trials (Pevernagie, 2021).

Future research should be aimed to resolve unanswered questions regarding OSA diagnosis and management (McNicholas et al., 2018) (Figure 4). New approaches to syndrome definition are required that consider different clinical OSA phenotypes. Certain comorbidities strongly associated with OSA could also be included in severity grading, if a causal association can be demonstrated. New diagnostic approaches are required that incorporate novel technologies to provide surrogates for sleep structure, to gauge exposure to systemic effects of OSA, and to identify specific biomarkers for disease classification. Indeed, both causality and susceptibility issues may be settled by finding markers that enhance specificity in the relationship between pathophysiological endotypes and clinical phenotypes. While useful markers can conceivably be derived from PSG (Gauld & Micoulaud-Franchi, 2021; Lechat et al., 2022; Lim et al., 2020; Pepin et al., 2018), it will be compulsory to adapt the conventional sleep diagnostic approach to techniques for ambulatory and multi-night assessment.

Eventually, these efforts will give way to a new framework for personalised treatment and precision medicine, not only by linking pathophysiology to phenotype, but also by providing markers that identify treatable traits suitable for targeted treatment (Bonsignore et al., 2017; Pevernagie, 2021; Pien et al., 2018). As OSA is a complex and heterogeneous disease that may overlap with other sleep disorders, a differentiated approach is mandatory. This observation underlines the importance of clinical sleep medicine, because of its integrative methodology and added value for managing person-specific sleep disorders.

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