Pathophysiology of Sleep Apnea

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Dempsey JA, Veasey SC, Morgan BJ, O’Donnell CP. Pathophysiology of Sleep Apnea. Physiol Rev 90: 47–112, 2010; doi:10.1152/physrev.00043.2008.—Sleep-induced apnea and disordered breathing refers to intermittent, cyclical cessations or reductions of airflow, with or without obstructions of the upper airway (OSA). In the presence of an anatomically compromised, collapsible airway, the sleep-induced loss of compensatory tonic input to the upper
airway dilator muscle motor neurons leads to collapse of the pharyngeal airway. In turn, the ability of the sleeping subject to compensate for this airway obstruction will determine the degree of cycling of these events. Several of the classic neurotransmitters and a growing list of neuromodulators have now been identified that contribute to neurochemical regulation of pharyngeal motor neuron activity and airway patency. Limited progress has been made in developing pharmacotherapies with acceptable specificity for the treatment of sleep-induced airway obstruction. We review three types of major long-term sequelae to severe OSA that have been assessed in humans through use of continuous positive airway pressure (CPAP) treatment and in animal models via long-term intermittent hypoxemia (IH): 1) cardiovascular. The evidence is strongest to support daytime systemic hypertension as a consequence of severe OSA, with less conclusive effects on pulmonary hypertension, stroke, coronary artery disease, and cardiac arrhythmias. The underlying mechanisms mediating hypertension include enhanced chemoreceptor sensitivity causing excessive daytime sympathetic vasoconstrictor activity, combined with overproduction of superoxide ion and inflammatory effects on resistance vessels. 2) Insulin sensitivity and homeostasis of glucose regulation are negatively impacted by both intermittent hypoxemia and sleep disruption, but whether these influences of OSA are sufficient, independent of obesity, to contribute significantly to the “metabolic syndrome” remains unsettled. 3) Neurocognitive effects include daytime sleepiness and impaired memory and concentration. These effects reflect hypoxic-induced “neural injury.” We discuss future research into understanding the pathophysiology of sleep apnea as a basis for uncovering newer forms of treatment of both the ventilatory disorder and its multiple sequelae.

I. INTRODUCTION

Sleep-disordered breathing refers to momentary, often cyclical, cessations in breathing rhythm (apneas) or momentary or sustained reductions in the breath amplitude (hypopneas), sufficient to cause significant arterial hypoxemia and hypercapnia. These apneas and hypopneas are specific to the sleeping state and are accompanied by 1) a compromised, often completely closed, extrathoracic upper airway (“obstructive” event); 2) a marked reduction or cessation of brain stem respiratory motor output (“central” event); and 3) a combination of central and obstructive events. These ventilatory inadequacies and their accompanying intermittent hypoxemia often lead to transient arousals from sleep and sleep state fragmentation throughout the night and cause compensatory responses of the autonomic nervous system. This phenomenon is now known to occur with varying degrees of severity in literally millions of people throughout the world. In our review we first provide a brief historical perspective of the problem and then detail the pathogenesis and selected long-term consequences of sleep apnea. We have not attempted to cover in depth all of the important research problems in what has become a large field of scientific endeavor. Instead, we refer the interested reader to recent reviews on the epidemiology of sleep-disordered breathing (446, 759), the influence of race and ethnicity (79, 334, 729), sleep apnea in children (543, 670) and the “sudden infant death” syndrome (304, 674), and the neurocognitive effects of sleep apnea (43, 204).

II. HISTORY OF SLEEP APNEA

We have only really begun to study and to understand sleep apnea over the past 40 years, even though there were strong hints of the widespread existence of this problem as early as the 19th Century. Observations of periodic breathing in sleep were first reported in the mid 1850s, and in the 1870s British physicians reported on several cases of obstructed apneas as “fruitless contractions of the inspiratory and expiratory muscles against glottic obstruction with accompanying cyanosis during sleep” (346). During the later half of the 19th century, several cases of obese persons with extreme daytime sleepiness were described (346) and labeled the “Pickwickian syndrome” after Charles Dickens Fat Boy Joe as described in the Pickwick Papers in 1837 (132). Periodic breathing was reported by British physician Hunter and by Irish physicians Cheyne and Stokes in heart failure patients in the early to mid 19th Century (101, 346, 649) and in otherwise healthy subjects sleeping in the hypoxia of high altitudes by the British Physiologists John Scott Haldane, C. G. Douglas, and Mabel Fitzgerald at the turn of the 20th Century (137).

It was not until the mid 1950s that a link between obesity and the control of breathing was fully appreciated as the Pickwickian syndrome was rediscovered. Daytime CO2 retention was observed in obese subjects with daytime sleepiness and without significant lung disease (55). Remarkably, any association with sleep disorders was not considered. In fact, the daytime sleepiness in these patients was ascribed to “CO2 poisoning” accompanying their respiratory failure. Indeed, respiratory physiologists and neurophysiologists studying the control of breathing in these times never considered the extrathoracic upper airway as an important factor in this control system, and we knew little about its neuromuscular regulation. Furthermore, even descriptions of sleep effects on ventilation and ventilatory stability in health were not reported until the comprehensive studies of Bulow in the early 1960s (78). Finally, by the mid 1960s, Gastaut et al. (188) recognized obstructive sleep apnea in obese subjects as intermittent airway obstruction with frequent arousals, thereby providing the first comprehensive
links between obesity, sleep-induced airway obstruction, sleep fragmentation, and daytime sleepiness. Following these key observations, research proceeded slowly with case reports of obstructive sleep apnea and the occasional use of chronic tracheostomy for treatment in the early 1970s (213, 375).

The findings of the mid to late 1970s through early 1980s provided a huge impetus to physiological research in this field of sleep and breathing, as highlighted by a series of reports of 1) sleep effects on brain stem respiratory neuronal activity in the unanesthetized cat (473, 474); 2) a neuromuscular reflex mechanism maintaining extrathoracic airway patency in the rabbit (74); 3) sleep effects on reflex control of breathing in the dog (514) and identification of a sensitive CO$_2$-induced apneic threshold in sleeping humans (621); 4) description of anatomical and neurophysiological determinants of upper airway occlusion in the sleeping human, which provided a unifying “balance of forces” concept of obstructive sleep apnea (OSA) pathogenesis (549); and 5) the landmark introduction of continuous nasal pressure (CPAP) application as the noninvasive treatment for obstructive sleep apnea (654). Early in the 1990s, simulation of OSA in rodents using cyclic hypoxia was shown to cause a gradual development of daytime hypertension, thereby initiating research into the long-term cardiovascular consequences of sleep apnea (171). At this same time OSA patients were shown to maintain their upper airway patency in wakefulness via a compensatory, augmented EMG activity of their airway dilator muscles (405), which extended an earlier report of more frequently occurring genioglossus EMG activity during wakefulness [and non-rapid eye movement (NREM) sleep] in OSA patients (657). Shortly thereafter, the first population study conducted using in-lab studies of sleep and breathing showed a significant prevalence of sleep apnea or sleep-disordered breathing in a middle-aged, nonclinical population (the Wisconsin Sleep Cohort), and these findings signaled a potentially significant and largely undiagnosed effect of sleep-disordered breathing on public health (758).

From the mid 1990s to the present, we have seen an explosion of basic, clinical, and population research directed toward the prevalence, causes, consequences, and treatment of this long-standing, although only recently appreciated, problem. Sleep apnea has attracted a myriad of researchers from diverse disciplines and clinical specialties. At the same time, sleep apnea as a serious, undefined clinical problem has also given birth to many commercial ventures for its diagnosis and treatment, including the building of literally hundreds of sleep medicine clinics throughout the western world with the majority of their business concerned with the diagnosis and treatment of sleep apnea. Finally, given the relatively high prevalence of this sleep-specific problem with potential carryover to daytime pathology, sleep apnea has provided great impetus to the growth of sleep medicine as a clinical and research specialty.

III. PATHOGENESIS OF SLEEP APNEA

A. Wakefulness Influences on Ventilatory Control

Remarkably, sleep apnea patients experience little or no problems with their breathing or airway patency while awake. In fact, the great majority of people with sleep apnea possess ventilatory control systems that are capable of precise regulation of their alveolar ventilation and arterial blood gases with extremely small variations from the norm throughout the waking hours. In addition, these healthy control systems, while awake, possess sufficiently sensitive feedback and feedforward controls to ensure precise coordination of chest wall and upper airway “respiratory” muscle recruitment so as to provide maximum airway diameter, low airway resistance and optimum lung volumes and respiratory muscle lengths, regardless of the ventilatory requirement.

To underscore the importance of the “waking stimuli” to breathe and to upper airway patency and to ventilatory control, consider the following qualitative influences of sleep on the control of breathing.

Electrical activity from medullary inspiratory neurons, EMG activity of diaphragm and abductor muscles of the upper airway in healthy humans and/or in cats, show reductions in amplitude upon the transition from awake to NREM sleep, usually accompanied by a mild to moderate hypoventilation (+2 to 8 mmHg Pa$_{CO_2}$) and two- to fivefold increases in upper airway resistance (128, 241, 369, 376). Sleep induces consistently greater proportional reductions in the EMG activity in the upper airway versus chest wall pump muscles (471).

A fast and highly variable breathing frequency is a hallmark of rapid eye movement (REM) sleep in mammals, even though postural muscles, including accessory respiratory muscles of the chest wall, are essentially atonic (513). So, an excitatory drive to breathe is common in REM coincident with increased diaphragmatic EMG activity and increased activity in many medullary respiratory neurons above those levels observed in NREM sleep or quiet wakefulness (467, 468, 470).

In cases of congenital central hypoventilation syndrome, the ventilatory response to imposed hypercapnia and to hypoxemia is absent; however, eupneic breathing rhythm is maintained while awake but lost completely in deep NREM or slow wave sleep (15, 160).

Pa$_{CO_2}$ can be lowered substantially (using mechanical ventilation) during wakefulness with little or no disruption of breathing pattern; however, in NREM sleep, very small transient reductions in Pa$_{CO_2}$ (even only to the waking level) result in significant apnea (239, 404, 621).
Partial ablation of the rat’s pre-Bötzinger complex, a major site of respiratory rhythm generation in the medulla, is without effect on breathing pattern or chemoresponsiveness in wakefulness but is accompanied by apnea and ataxic breathing patterns in NREM and REM sleep (401). Furthermore, focal acidification of the retrotrapezoid nucleus, a major site of medullary chemoreception, produces a significant ventilatory response in wakefulness, with no response in sleep (353).

Added resistive or elastic loads to the airway prompt immediate and highly variable increases in the drive to breathe in the waking state, which prevent hypoventilation; however, in sleep, these mechanical loads are not accompanied by an immediate compensatory increase in the drive to breathe, and hypoventilation ensues until chemoreceptor stimuli increase (240, 261, 728).

Mechanical or chemical stimulation of the larynx or intrapulmonary airways causes cough in wakefulness but not in NREM or REM sleep (655), implying that acts requiring complex coordination of glottal, intercostal, diaphragm, abdominal, and tracheobronchial muscles require participation of supramedullary structures that are activated and synchronized while awake but not in NREM sleep.

Studies in chronically instrumented cats and rats have provided findings which demonstrate that the wakefulness stimuli to breathe include tonic excitatory inputs from the reticular formation, brain stem amnergic systems, and hypothalamic orexin-containing neurons (469). In NREM sleep, decrements occur in these excitatory inputs, and in REM sleep, there are both tonic excitatory inputs and phasic inhibitory inputs that account for irregularities in breathing pattern as well as the loss of excitation which contributes to hypotonia of the muscles of the upper airway (163, 259, 467–469, 472, 553, 641). Details concerning sleep effects on the neurochemical control of breathing and airway patency are provided in section IV.

In most healthy human subjects of any age, sleeping at low altitudes, the loss of these wakefulness influences on neurochemical control of breathing and airway patency is of minor physiological consequence. Mild CO₂ retention and respiratory acidosis and reductions in alveolar Po₂ are not accompanied by significant arterial O₂ desaturation or a compromised systemic O₂ transport (because of the high affinity of Hb for O₂). Furthermore, while loss of upper airway muscle tonic activity results in a doubling or even sometimes a quadrupling of upper airway resistance and intrathoracic pressure swings in sleep in many healthy humans, there are usually only small, inconsequential effects on pulmonary gas exchange, sleep state continuity, autonomic regulation, or ventricular function. However, for many otherwise healthy subjects in whom airway patency is already anatomically compromised in wakefulness and/or whose ventilatory control systems are inappropriately driven by chemical stimuli, simply loss of wakefulness inputs to the control of the upper airway and chest wall muscle motor neurons produces serious, short- and long-term consequences to homeostasis and to health. We shall now discuss the mechanisms contributing to the complex pathogenesis of sleep-disordered breathing.

B. Defining Sleep-Disordered Breathing Events and a Roadmap for Pathogenesis

Sleep-disordered breathing leading to repeated bouts of ventilatory overshoots and undershoots and accompanying swings in arterial blood gases and intrathoracic pressure takes on many forms. Commonly, sleep-disordered breathing is divided into so-called “central” events, denoting an absence or marked reduction in central respiratory motor output to respiratory pump muscles, or “obstructive” events, which are comprised of respiratory efforts against a closed upper airway. However, as we discuss below, most cyclical sleep-disordered breathing events are driven by anomalies in both anatomical and neurochemical control of upper airway and/or chest wall respiratory musculature. Four patterns of sleep-disordered breathing are illustrated in Figure 1, including the waxing and waning of ventilatory responses in the severe heart failure patient (see Fig. 1A), the “cluster” periodic breathing of healthy sea-level natives during sleep in the hypoxia of high altitude (Fig. 1B), and the obstructive and “central” apneas coexisting in an OSA patient during sleep (Fig. 1C). These examples represent the extremes of a broad continuum of sleep-disordered breathing that also includes airway narrowing, rather than complete airway obstruction and periods of transient hyperventilation or “hypopnea” rather than complete apnea.

Population-based cross-sectional and longitudinal studies have specified the dominant risk factors for OSA in the general (nonclinical) population to include excess body weight as the dominant contributor, followed by male gender, and to significant but lesser extents, cranial facial structures and aging (129, 488, 501, 562, 757–759). We will concentrate on the more common affliction of OSA. In short, the process is initiated because the wakeful state provides compensatory neuronal activation of dilator muscles in an anatomically compromised collapsible pharynx; accordingly, when this activation is lost at sleep onset, the airway narrows and/or collapses. However, the tendency to result (or not to result) in repeated cyclical apneas is the end product of multiple compensatory processes that vary markedly among and within individuals. Concepts have continued to evolve as we learn more about the neurophysiological mechanisms governing control of respiratory rhythm and its coupling with upper airway control and states of consciousness and applying these principles to human patients during sleep.

We discuss OSA pathogenesis in three steps, as outlined in Figure 2. First we detail the varied structural and...
functional determinants of an anatomical predisposition for airway closure, an absolutely essential component for OSA. The second essential component is sleep. This section emphasizes the effects of the sleeping state on mechanisms underlying both obstructive and central apnea and ventilatory instability. Finally, we attempt to integrate anatomical deficits with mechanisms underlying central neurochemical control of breathing stability and compensatory neuromuscular control of upper airway caliber, to explain the cyclical, repetitive nature of OSA.

C. Anatomical Determinants of Upper Airway Caliber in OSA

1. Unique anatomy of the human airway

The upper airway is a complex structure required to perform deglutition, vocalization, and respiration. In the human, this structure must also perform tightly controlled and complex motor behaviors required for speech. Upper airway obstruction in sleep is most prevalent in the human in part because the hyoid bone, a key anchoring site for pharyngeal dilator muscles, is not rigidly attached to skeletal structures. In other mammals, the hyoid bone is attached to the styloid processes of the skull (425, 756). Thus the human pharynx has no rigid support except at its extreme upper and lower ends where it is anchored to bone (upper) and cartilage (larynx); therefore, pharyngeal cross-sectional area will vary with lumen pressure (271). Humans depend critically on the coordinated actions and interactions of over 20 skeletal muscles that dilate and stent open the oropharynx (see sect. IV C).

Beyond the hyoid arch, Lieberman et al. (361, 362) and Davidson (117) also point to the anatomical changes in the adult human upper airway during the evolutionary development of speech as a potential major contributor to OSA. Specifically, the gradual decent of the larynx to a position greatly inferior to the oropharynx separated the soft palate from the epiglottis. The creation of this “supralaryngeal vocal tract,” like the tube of a clarinet, “filters” the sound produced by the larynx and in turn speech is produced via the changing position of the pharynx, tongue, and lips. The downside is a relatively shortened, compacted face and greatly narrowed oropharynx in

FIG. 1. A: periodic (Cheyne-Stokes) breathing in chronic heart failure in non-REM sleep. Note the gradual crescendo and decrescendo of tidal volume ($V_t$) and esophageal pressure ($P_{es}$), the intermittent hypoxemia ($SaO_2$), and the subtle changes in EEG amplitude attending the termination of each periodic breathing cycle. Periodic cycles of apnea plus hyperpnea are fairly uniform and are each 50–60 s in duration. [From Tkacova et al. (681).] B: periodic “cluster-type” breathing in non-REM sleep in a healthy sea-level native during the initial night at 4,300 m altitude. Tidal volume is estimated from expansion of the ribcage (RC) and abdomen (Abd) using inductance plethysmography. Note the abrupt increase in $V_t$ (to 1.5–2.5 times control, steady-state values) at the end of each apneic period. Each periodic cycle is 20–25 s in length. Also note the mild levels of arterial hypocapnia and alkaline pH determined from blood sampling over several periodic cycles. [From Berssenbrugge et al. (53).] C: cyclical “mixed,” i.e., central followed by obstructed apneas, causing intermittent hypoxemia during non-REM sleep. The cessation of airflow denotes the onset of apnea. The absence of cyclical changes in esophageal pressure over the initial 8–10 s of the apnea demonstrates that this initial phase of the apnea is due to the absence of “central” respiratory motor output and inspiratory muscle contractions. Over the latter half of the apnea, flow is still absent but progressive and cyclical increments occur in the negativity of esophageal pressure, indicating increasing inspiratory efforts against a closed airway in response to rising asphyxic chemoreceptor stimuli. The arrows shown at the termination of each apneic period indicate periods of transient cortical arousal.
which the tongue encroaches significantly on the available space.

2. Sites of airway collapse

Studies using nasal pharyngoscopy, computer tomography and magnetic resonance imaging, or pharyngeal pressure monitoring have shown that one or more sites within the oral pharyngeal region are usually where closure occurs in most subjects with OSA, and this region is also smaller in OSA patients versus controls even during wakefulness (see Fig. 3A) (253, 426, 597, 599). Although the retropalatal region of the oropharynx is the most common site of collapse (see Fig. 3B), airway narrowing is a dynamic process, varying markedly among and within subjects and often includes the retroglossal and hypopharyngeal areas (255, 430, 444). For example, Watanabee et al. (710) have shown that airway closure in obese OSA subjects occurred primarily at the velopharynx, whereas in nonobese OSA patients with a retracted mandible, the closure occurred at both the velo- and oropharynx.

3. Soft tissue and bony structure abnormalities

The recent use of quantitative imaging techniques has allowed advances that reveal important differences in both craniofacial and upper airway soft tissue structures in the OSA patient. The reduced size of cranial bony structures in the OSA patient include a reduced mandibular body length, inferior positioned hyoid bone, and retro position of the maxilla, all of which compromise the pharyngeal airspace (21, 556, 557). Airway length, from the top of the hard palate to the base of the epiglottis, is also increased in OSA patients, perhaps reflecting the increased proportion of collapsible airway exposed to collapsing pressures (385, 479). As expected, these craniofacial dimensions are primarily inherited, as the relatives of OSA patients demonstrated retracted and short mandibles and inferiorly placed hyoid bones, longer soft palates, wider uvulas, and higher narrower hard palates than matched controls (214, 396).

Enlargement of soft tissue structures both within and surrounding the airway contributes significantly to pha-
Ryngeal airway narrowing in most cases of OSA. An enlarged soft palate and tongue would encroach on airway diameter in the anterior-posterior plane (107, 254), while the thickened pharyngeal walls would encroach in the lateral plane. Volumetric time overlapped magnetic resonance imaging (MRI) or computer tomography (CT) images strongly implicate the thickness of the lateral pharyngeal walls as a major site of airway compromise, as the airway is narrowed primarily in the lateral dimension in the majority of OSA patients (505, 597). Furthermore, treatment with CPAP, weight loss, or mandibular advancement all show increases in the lateral pharyngeal dimensions (561, 595, 597).

There are many potential causes of lateral wall thickening in OSA patients. First, as shown in both humans and rodent models, obesity is a major contributor to airway compression through increased area and volume of pharyngeal fat deposits (69, 253, 606, 610). This excess fat deposition has also been observed under the mandible and within the tongue, soft palate, or uvula (645). Obesity also gives rise to excess fat-free muscular tissue, thereby increasing the size of many upper airway structures (65, 253, 600, 606) and compressing the lateral airway walls. In children with OSA, tonsillar and adenoid hypertrophy form the major anatomical contributors to airway narrowing (388).

4. Obesity and lung volume

Obesity also contributes indirectly to upper airway narrowing, especially in the hypotonic airway present during sleep, because lung volumes are markedly reduced by a combination of increased abdominal fat mass and the recumbent posture. In turn, the reduced lung volume...
reduces the “tug” on the trachea induced by the traction exerted via mediastinal structures by negative intrathoracic pressures and by the diaphragm descent, thereby further increasing the thickness of the lateral pharyngeal walls and narrowing the airway.

The highly sensitive “traction” effect of changes in lung volume on upper airway patency and airway resistance was clearly demonstrated in anesthetized animals by Van de Graaff who surgically disconnected the mechanical linkage between chest wall and upper airway by severing all cervical structures anterior and anterolateral to the spine (690). While intact, upper airway resistance ($R_{ua}$) fell during inspiration, whereas following removal of the mediastinal-tracheal linkages $R_{ua}$ was increased and no longer underwent respiratory modulation. Lung volume effects on $R_{ua}$ are also mediated in part via pharyngeal dilator muscle activity; however, these lung volume effects persist even following denervation of airway muscles in the dog (690) or during muscle paralysis in the human (663). Also with humans sleeping in an iron-lung respirator, variations in box pressure surrounding the chest and abdomen produced small changes in end-expiratory lung volume (EELV), which in turn produced highly sensitive effects on upper airway patency and resistance (45, 235) and on airway closing pressures (see sect. II.D), even in the paralyzed patient (663). The greater the upper airway collapsibility and airway resistance in these sleeping subjects, the more the airway resistance was reduced as EELV was increased (605, 607, 644). Furthermore, based on endoscopic imaging studies, lung volume effects on airway collapsibility were shown to be more pronounced at the level of the velopharynx versus oropharynx and in obese versus nonobese subjects (663). This sensitive, purely mechanical effect of lung volume changes on passive control of upper airway patency and resistance has not been fully appreciated to date; indeed, this effect may explain at least some of the effect of obesity, sleep itself, and even CPAP treatment on upper airway collapsibility (602) and improve apnea hypopnea index (AHI) by 20–30% (283, 305, 428).

5. Airway edema and surface tension

Accumulation of even relatively small amounts (~100–200 ml) of edematous fluid enlarges upper airway soft tissue structures in OSA patients and snorers, especially in the soft palate which may be tugged caudally and constricted during apneas (596). Local vascular engorgement may also enlarge soft tissues in the upper airway (709). Furthermore, cephalad displacement of fluid from the lower extremities to the upper airway upon assuming recumbency has been recently documented and associated with sleep-disordered breathing (105, 544, 653).

Surface tension of the liquid lining the mucosa affects collapsibility of the upper airway in the same way as it has been well documented in the lung’s airways. A higher surface tension in the upper airway wall of OSA patients has been reported using a method that quantifies surface tension as the force required to separate two surfaces bridged by a droplet of the liquid under study (305, 306). Furthermore, in limited studies, surfactant therapy in OSA patients was shown to significantly reduce airway collapsibility (602) and improve apnea hypopnea index (AHI) by 20–30% (283, 305, 428).

6. Obesity, leptin, and inflammation

Central, or visceral, obesity is associated with the greatest risk for OSA (611). This suggests that factors other than pure mechanical load may contribute to the pathogenesis of respiratory disturbances during sleep. The concept is now emerging that visceral fat deposits, which represent a rich source of humoral mediators and inflammatory cytokines, can impact on neural pathways associated with respiratory control (601). Perhaps the most well-studied adipocyte-derived factor affecting respiratory control is leptin, which was initially determined to have a primary role of binding to receptors in the hypothalamus to reduce satiety and increase metabolism (178). Leptin can also act as a respiratory stimulant, and impairment of the leptin signaling pathway, as occurs in leptin-resistant or leptin-deficient states of obesity, causes respiratory depression in mice (453) and is associated with obesity hypoventilation syndrome in humans (515). Even though obesity and OSA are associated with elevated circulating levels of leptin, if centers in the brain impacting on respiratory control act in a similar leptin-resistant manner to hypothalamic regions controlling appetite and metabolism, then impaired leptin signaling in the CNS may contribute to respiratory depression as predicted in murine studies.

In addition to respiratory control, animal studies show leptin is also critical in lung development and affects the distribution of muscle fiber types in the diaphragm (667). However, as yet there is no direct evidence that impaired leptin signaling can impact on the control of respiratory muscles of the upper airway, although it may play a role in nocturnal hypoventilation, particularly in REM sleep where respiration is markedly depressed in leptin-deficient mice (453). Visceral adipose tissue releases many other humoral factors including classical proinflammatory cytokines such as tumor necrosis factor-α (TNF-α) and interleukin (IL)-6 that are elevated in OSA and can be reduced with CPAP therapy (409, 749). These and other proinflammatory cytokines may impact on sleep (613) or potentially contribute to the local in-
flammary response reported in the upper airway tissues of OSA patients (64), but there is little evidence of an effect on respiratory control. Overall, the concept that adipocyte-derived circulating factors can impact on respiratory control of the upper airway or act on the upper airway directly to contribute to the pathogenesis of OSA is intriguing, but currently lacking clear supporting data.

Finally, in our emphasis on the importance of obesity both independently and in its interactions with gender, age, lung volume, and cranial facial dimensions, we need to acknowledge the accumulation of evidence from genome-wide linkage studies of OSA phenotypes (481, 482) pointing to a strong heritability of both AHI (~33%) and body mass index (BMI) (>50%), with about one-half of the genetic determinants of AHI being obesity related and one-half being independent of obesity (489, 760). A heritable predisposition to this disorder is also suggested by its higher prevalence in males versus females and in African ancestors and East Asians compared with Western populations (269, 543, 758). The wide array of candidate genes that might link genetic mechanisms of obesity with sleep apnea are under investigation and have been recently reviewed (489, 760).

D. Mechanical Determinants of Upper Airway Patency

Mechanical determinants of airway caliber of the human pharynx in sleep are similar to those regulating caliber of any collapsible tube (598). Other well-known biological examples in respiratory physiology include intrathoracic airway collapse upon forced exhalation (531), collapse of pulmonary capillaries in the lung apex (718), and collapse of alae nase at high inspiratory flow rates (70). A Starling resistor model developed by Schwartz and colleagues (603, 628) consists of a collapsible tube with a sealed box interposed between two rigid segments (195) (see Fig. 4). The critical closing pressure ($P_{\text{crit}}$) of the passive airway is defined as the pressure inside the airway ($P_{\text{in}}$) at which the airway collapses. The pressure gradient during airflow through the system is defined by $P_{\text{upstream}} - P_{\text{downstream}}$. Therefore, with increasing $P_{\text{crit}}$, as the differential between $P_{\text{upstream}}$ and $P_{\text{crit}}$ decreases, inspiratory airflow limitation will eventually develop, and when the $P_{\text{us}}$ falls below $P_{\text{crit}}$, complete airway occlusion occurs. Effective therapy for sleep apnea requires that the $P_{\text{us}}$ to $P_{\text{crit}}$ pressure differential be widened, and this can be accomplished by either 1) an increase in $P_{\text{us}}$ with appropriate amounts of CPAP applied at the airway opening, or 2) by decreasing $P_{\text{crit}}$ via either reducing the collapsing pressures on the airway (e.g., weight loss or alteration of cranial-facial anatomy or increasing lung volume) or by augmenting “active” neuromuscular control of airway tone (see sect. mE) (195, 492).

Airway $P_{\text{crit}}$ due solely to mechanical properties of the airway and its surrounding tissue, termed “passive $P_{\text{crit}}$ ”, has been assessed during sleep and under conditions of complete muscle atonicity in paralyzed, anesthetized patients through the use of pressure control systems connected to nasal masks that are capable of manipulating airway pressure in a stepwise fashion across a wide range (~20 cmH2O) (195, 271, 272, 492). In general, these measures in sleeping or paralyzed humans have shown that passive $P_{\text{crit}}$ is in the range of < −10 cmH2O in normal subjects with low airway resistance and minimal CO2 retention during NREM sleep, from −10 to −5 cmH2O in snorers, −5 to 0 cmH2O in those with sufficiently high airway resistance to induce airflow limitation and transient hypopneas, and >0 cmH2O in patients with apneas with complete airway obstruction.

Passive $P_{\text{crit}}$ is significantly associated with the two greatest risk factors for OSA, namely, obesity (see sect. mC) and gender. Recent studies using substantial numbers of males and females matched for BMI and age have documented a substantially greater 2–3 cmH2O elevation in passive $P_{\text{crit}}$ (during NREM sleep) in men versus premenopausal women (285, 307). This gender difference in passive airway collapsibility may be related to the longer pharyngeal airway length and in the mass of soft tissue contained in the soft palate and tongue in males (385, 722). In turn, this difference in pharyngeal soft tissue.

**FIG. 4.** Starling resistor model of obstructive sleep apnea. In the Starling resistor model, the collapsible segment of the tube is bound by an upstream and downstream segment with corresponding upstream pressure ($P_{\text{us}}$), downstream pressure ($P_{\text{ds}}$), and upstream resistance and downstream resistance. Airway occlusion occurs when the surrounding tissue pressure ($P_{\text{tu}}$), (comprised of pharyngeal muscles and pharyngeal and submucosal fat, mucosal edema, etc.; see sect. mC), becomes greater than the intraluminal pressure ($P_{\text{in}}$), resulting in a transmural pressure of zero. In this model of the upper airway, $P_{\text{in}}$ is atmospheric at the airway opening, and $P_{\text{tu}}$ is the tracheal pressure. The critical closing pressure of the collapsible airway ($P_{\text{us}}$) is represented by $P_{\text{in}}$. The Starling resistor model, the collapsible segment of the tube is bound by an upstream and downstream segment with corresponding upstream pressure ($P_{\text{us}}$), downstream pressure ($P_{\text{ds}}$), and upstream resistance and downstream resistance. Airway occlusion occurs when the surrounding tissue pressure ($P_{\text{tu}}$), (comprised of pharyngeal muscles and pharyngeal and submucosal fat, mucosal edema, etc.; see sect. mC), becomes greater than the intraluminal pressure ($P_{\text{in}}$), resulting in a transmural pressure of zero. In this model of the upper airway, $P_{\text{in}}$ is atmospheric at the airway opening, and $P_{\text{tu}}$ is the tracheal pressure. The critical closing pressure of the collapsible airway ($P_{\text{us}}$) is represented by $P_{\text{in}}$. When the $P_{\text{tu}}$ is significantly lower than $P_{\text{us}}$ and $P_{\text{ds}}$, flow through the tube occurs. When $P_{\text{us}}$ falls during inspiration below $P_{\text{crit}}$, inspiratory airflow limitation occurs and is independent of further decreases in $P_{\text{ds}}$. Under this condition, the pharynx is in a state of partial collapse, and maximal inspiratory airflow varies linearly as a function of the difference between $P_{\text{us}}$ and $P_{\text{crit}}$. Finally, when $P_{\text{us}}$ falls below $P_{\text{crit}}$, the upper airway is completely occluded. [Adapted from Gold and Schwartz (195).]
deposition may be secondary to the tendency toward fat deposition in the upper body and trunk in males versus lower body and extremities in females (407).

E. Neuromuscular Control of Upper Airway Dynamics in Sleep

Clearly then the effects of airway anatomy on airway collapsing pressure in a hypotonic airway are a critical determinant of obstructive apnea. However, several lines of evidence also support neuromuscular factors as significant determinants of airway collapsibility in sleep. First, tonic and phasic EMG activity of pharyngeal airway dilator muscles (genioglossis and tensor palatine) are progressively reduced from wakefulness to NREM to REM sleep and further inhibited coincident with the “phasic” eye movement events in REM. This powerful effect of state has been adequately documented in tracheostomized animal models (374) and recently has been demonstrated in OSA patients in whom the potentially confounding, compensatory responses to sleep-induced changes in upper airway resistance, negative pressure, $P_{\text{CO}_2}$, and respiratory motor output were controlled through the use of either CPAP (148) or positive pressure controlled mechanical ventilation (369). These state effects on the neuromuscular control of the upper airway likely explain, along with reductions in lung volume (see sect. III.C), why $P_{\text{crit}}$ is never positive in the waking state, even in OSA patients. Furthermore, genioglossus EMG activity is abnormally high in awake OSA patients (405, 657), and its sleep-induced decrement may be viewed as playing a “permissive” role in explaining (or unmasking) closure of an already anatomically compromised upper airway (see sect. IV.D for neurochemical mechanisms underlying these state effects).

Second, neuromuscular factors also play a significant role in the dynamic breath-to-breath and intrabreath regulation of upper airway caliber, through changes in proprioceptive and chemoreceptor feedback. During inspiration, the passive pharynx narrows as intraluminal pressure is progressively reduced because of energy lost in overcoming frictional airway resistance and increases in flow velocity secondary to the Bernoulli effect operating in a reduced lumen size (598). This collapsing effect of a reduced luminal pressure is opposed during inspiration by a reduction in dynamic compliance, i.e., collapsibility, of the airway achieved via reflex activation of pharyngeal dilator muscles. A manifestation of this dilator muscle recruitment is reflected in the markedly higher active $P_{\text{crit}}$ obtained in OSA patients when they breathe through a tracheostomy versus nasal breathing, pointing to the significant activation of upper airway muscles during inspiration through the intact upper airway (384, 591). In turn, the reflex activation occurs in response to negative pressure airway mechanoreceptors located principally in the larynx and to a lesser extent in the superficial layers of the pharyngeal wall, with their afferent projections located in the superior laryngeal nerve, and also in glossopharyngeal and trigeminal nerves (250, 260, 323, 394, 395, 579, 591, 691). Large changes in negative pressure in the isolated upper airway trigger a dual protective reflex, which restores airway patency by both activating airway dilators (to reduce airway compliance) while inhibiting diaphragm EMG activity (which minimizes intraluminal negative pressure) (224, 394). Vagally mediated feedback influences on laryngeal, tongue, and hyoid muscle via pulmonary stretch receptors also protects against airway collapse as the rate of lung inflation is slowed in the face of increased airway resistance, thereby reflexly activating upper airway motor neurons (323). Finally, chemoreceptor influences also have substantial effects on upper airway muscle recruitment, and in the case of $CO_2$, upper airway motor neurons relative to phrenic motor neurons have been shown to have a substantially higher threshold for inhibition (via hypocapnia) and activation (via hypercapnia) (466, 711).

Dynamic imaging of upper airway caliber as well as breath by breath analysis of airway mechanics during sleep shows that narrowing/closure may occur at endexpiration or during inspiration (25, 426, 574, 575, 658), each of these airway occurrences suggesting quite different mechanisms precipitating collapse. End-expiratory occlusion occurs without the need for generating an inspiratory effort or negative intraluminal pressure and may reflect that at end expiration the airway is no longer held open by phasic inspiratory activation of upper airway dilator muscles or by positive intraluminal pressure (25, 426, 595, 596). On the other hand, closure during inspiration points to an imbalance between the generation of upper airway muscle dilating forces versus an excessive intraluminal negative pressure generated by inspiratory chest wall muscles (549, 598, 658). Circumstances that might favor expiratory over inspiratory phase airway closure have not been thoroughly investigated, although limited data in the anesthetized obese mouse (69) and British bulldog (696) suggest that with anatomically compromised airways, expiratory phase narrowing and inspiratory phase (active) dilation are common in obesity, and the opposite effects occur in the airways of normal, nonobese control animals. Most evidence appears to support a passive closure of the upper airway during expiration as the dominant occurrence in OSA.

In summary, the evidence to date supports important roles for both anatomical and neural control of dilator muscles to the regulation of upper airway caliber in the sleeping human. The relative contributions of these factors will vary widely among and within individuals with, for example, patterns of fat deposition on the one hand and neurochemical sensitivity for dilator muscle recruitment on the other. Important indirect influences on air-
way caliber may also occur through instabilities in respiratory motor output, as we now discuss below.

**F. Instability of Central Respiratory Motor Output and Breathing Pattern in Sleep**

1. **Unmasking a sensitive apneic threshold**

Central apneas and instabilities in humans occur primarily in NREM sleep because of the critical dependence of the ventilatory control system on chemical stimuli, principally Pa\textsubscript{CO\textsubscript{2}}, in this state. Thus when normal subjects are mechanically ventilated in NREM sleep, transient increases in tidal volume and small reductions in Pa\textsubscript{CO\textsubscript{2}} only to waking levels (−3 to 5 mmHg) are sufficient to induce central apnea (239, 404, 621, 677) (see Fig. 5). A similar sensitive hypocapnic-induced apneic threshold can be demonstrated in sleeping, tracheostomized OSA patients (262) or dogs (106) by causing brief airway occlusions which in turn cause chemoreceptor (and often arousal)-driven transient ventilatory overshoots upon termination of the occlusion with subsequent central apneas (upon resumption of sleep). So apnea occurs whether the transient hyperventilation and hypocapnia are produced via “active” or “passive” means. Normally, in wakefulness, a transient increase in central respiratory motor output and hyperventilation are not followed by apneas, because a centrally mediated short-term potentiation of central respiratory motor output lingers following cessation of the ventilatory stimulus and ventilation returns gradually to its control eupneic levels (22, 152). However, in sleep, this stabilizing influence is apparently overridden by a transient reduction in the CO\textsubscript{2} drive to breathe, i.e., a sensitive apneic threshold is unmasked. Further evidence for a pivotal role for hypocapnia is the consistent evidence that the prevention of central apnea and periodic breathing during sleep in heart failure (734) or in hypoxic environments (53) is readily achieved by the addition of even very small amounts of inspired P\textsubscript{CO\textsubscript{2}}, i.e., just sufficient to prevent the occurrence of transient hypocapnia, regardless of the magnitude of any ventilatory overshoot.

Of course to reach the apneic threshold we need a source of transient ventilatory overshoots. Transient arousals from sleep provide a common source of this transient extra-drive to breathe, and the accompanying reductions in upper airway resistance permit a greater hyperventilatory manifestation of this increased drive (252, 743, 753). Arousals are especially effective in causing ventilatory overshoots when combined with increasing chemoreceptor drives from the preceding ventilatory undershoot (106, 262). Furthermore, the P\textsubscript{aCO\textsubscript{2}} required to terminate apnea is estimated to be 1–4 mmHg higher than the threshold P\textsubscript{aCO\textsubscript{2}} needed to initiate the apnea, reflecting a so-called control system “inertia” which delays the resumption of breathing rhythm (347, 581). The resultant apnea prolongation presents an enhanced chemoreceptor stimulus to ventilatory overshoot at apnea termination. While the hypocapnic-induced apnea threshold is highly sensitive and reproducible in NREM sleep, there is no apparent threshold in phasic REM sleep even with marked reductions in P\textsubscript{ETCO\textsubscript{2}} produced by either mechanical ventilation or by ventilatory overshoots in response to experimental airway occlusion (731). The central apneas and the periodic breathing accompanying heart failure or high-altitude hypoxia are also rarely present in REM sleep (53, 221). Perhaps, analogous to the wakeful state, the

**Fig. 5.** Effects of spontaneous central apnea on upper airway patency during NREM sleep. Fiber optic nasopharyngoscopy was used to determine airway dimensions at the level of the velo- or oropharynx. The initiation of central apnea is identified by the open inverted arrow, with the cessation of both airflow and oscillation of esophageal pressure (P\textsubscript{es}). Complete airway occlusion occurred −10 s following the onset of central apnea and before an inspiratory effort occurred, as noted by the constant P\textsubscript{es}. Central apnea continued and the airway remained closed for 35 s, showing partial return of airflow with resumption of inspiratory effort and then complete airway patency on arousal from sleep with an accompanying ventilatory overshoot. [From Badr et al. (25).]
erratic, sporadic increases in central inspiratory neural drive during REM (467) override hypocapnic inhibition.

Although hypocapnia is required during the ventilatory overshoot to cause subsequent apnea, lung stretch (732) and/or increases in systemic blood pressure and baroreceptor stimulation (582, 726) accompanying the ventilatory overshoot may also contribute to the ventilatory depression following the overshoot. Several reflex effects of airway negative pressure-sensitive mechanoreceptors on ventilatory control have also been demonstrated in the sleeping canine with an isolated upper airway. These include 1) the inhibitory effects of flow through the upper airway, or lung inflation on the rate of rise of diaphragmatic EMG activity; and 2) the apnea caused by either negative pressure pulses or low-pressure high-frequency pressure oscillations (akin to human snoring) applied to the airway during early expiration (147, 224, 520).

2. Sites of chemoreception causing apnea and instability

At what chemoreceptor site is transient hypocapnia acting in causing central apnea and periodic breathing? Carotid body denervation studies in the sleeping animal showed that neither apnea nor periodic breathing could be elicited even when substantial levels of transient hypocapnia were produced via mechanical ventilation (437). Similarly, using an isolated perfused carotid body preparation in the neurally intact sleeping canine (624, 626), central chemoreceptor hypocapnia (alone) was shown to produce little prolongation of expiratory time ($T_E$) despite marked reductions in $P_{aCO_2}$ (8 to 15 mmHg), whereas transient hypocapnia of only 3–5 mmHg produced apnea and periodic breathing when both chemoreceptors were able to sense the hypocapnia.

These data are strongly supportive of peripheral over central chemoreceptor in eliciting hypocapnic-induced apneas. On the other hand, we also know that perfusion of the isolated carotid chemoreceptor alone with severely hypocapnic (or hyperoxic) blood will reduce tidal volume (VT) but not cause apnea (627). Furthermore, specific increases in brain extracellular fluid (ECF) $[H^+]$ cause substantial increases in ventilation (161, 441, 626, 666). Accordingly, these apparent contradictions point to a new perspective put forward primarily by Guyenet (215) which emphasizes the potential importance of interdependence between peripheral and central chemoreceptors in ventilatory control.

Neuroanatomical evidence in the rodent model shows that Phox 2b gene expression delineates an uninterrupted chain of neurons in a circuit which includes carotid bodies and their afferent projections (116), chemoreceptor projections of the nucleus tractus solitarius (NTS) to the ventral lateral medulla (VLM) (666), and to $CO_2$ sensitive chemoreceptor neurons in the retrotrapezoid nucleus (RTN) (650). Functionally, an important interdependence between the various chemo- and mechanoreceptor afferents in the respiratory control system has also been demonstrated in reduced preparations by 1) the marked effect of systemic hypoxia increasing the activity of central $CO_2$-sensitive neurons in the RTN, an effect which was subsequently eliminated via carotid body denervation (650, 666); 2) the powerful hyperadditive effects of varying levels of $P_{CO_2}$ in the isolated perfused medulla on the ventilatory response to specific carotid body stimulation in the decerebrate vagotomized rat (121); and 3) the effect of vagal inhibition via lung stretch on carotid chemoreceptor (30) and medullary chemoreceptor responsiveness (215).

Further advances in our understanding of the chemical control of breathing requires that we no longer view the peripheral and central chemosensors as “stand alone” receptors, responding only to changes in the ionic composition of their immediate environment. Rather, we need to determine whether these proposed interdependencies are additive, hypoadditive, or hyperadditive in their influences on the final respiratory motor output (to both upper airway and chest wall) under conditions of transient and steady-state changes in systemic chemical stimuli and during wakefulness and sleep in neurally intact, fully responsive animal models.

3. Variations in susceptibility to central apnea and ventilatory instability

The occurrence of central apneas with repeated cyclic periods of over- and underventilation during sleep varies markedly depending on the gains(s) of the respiratory control system and the stability of the sleeping state. The tendency toward instability depends on the respiratory control system’s “loop gain,” an engineering term which defines the “gain” of the negative-feedback loop which regulates ventilation in response to a ventilatory disturbance (302). For example (see Fig. 6), if the magnitude of the increase in ventilation is greater than or equal to the magnitude of the preceding apnea or hypopnea, i.e., a high loop gain, then the system is highly unstable and will fluctuate between under- and overventilation (715).

Two types of control system gain, controller gain and plant gain, are major determinants of loop gain and therefore ventilatory stability (99, 300, 302). We illustrate the effects of changing each of these gains on $CO_2$ responsiveness above and below eupnea in Figure 7, which in turn will determine the tendency of ventilatory drive to overshoot in response to a rising chemoreceptor stimulus or to be overly depressed in response to the ensuing hypocapnia (127). For example, Figure 7A illustrates the effect of changing the background drive to breathe which will displace the eupneic $P_{aCO_2}$ along the isometabolic
line defining the hyperbolic relationship of $\text{PaCO}_2$ to alveolar ventilation. This hyperventilation, per se, protects against apnea and ventilatory instability by requiring a larger additional transient hyperventilation and hypocapnia to reach the apneic threshold (i.e., decreased plant gain), whereas a reduced drive and hypoventilation make one highly susceptible to apnea, requiring only very small further transient ventilatory overshoots (increased plant gain) (436). The other means of changing the magnitude of the $\text{CO}_2$ reserve below eupnea is to change the slope of the change in ventilation above and below eupnea, respectively, in response to induced hyper- or hypocapnia (see Fig. 7B, bottom). For example, an increased $\text{CO}_2$ response slope above and below eupnea (increased controller gain) has been observed in hypoxic humans and dogs and in chronic heart failure patients who experience periodic breathing in sleep (436, 737, 739, 742). This increased slope of response to changes in $\text{PaCO}_2$ results in a reduction of the $\text{CO}_2$ reserve and an increased susceptibility to apnea and periodicity despite the background hyperventilation, reduced eupneic $\text{PaCO}_2$ and plant gain (127). Thus the magnitude of loop gain and of the $\text{CO}_2$ reserve and therefore the propensity for ventilatory instability in sleep is determined by the net effects of any changes or abnormalities in controller gain vs. plant gain.

4. Cerebral blood flow and ventilatory instability

Cerebral vascular responsiveness to $\text{CO}_2$ is an important protector of brain extracellular fluid $\text{PCO}_2$ and $[\text{H}^+]$ by means of regulating cerebral blood flow (CBF) and the arterial to brain $\text{PCO}_2$ difference (587–589). For example, any reduction in cerebral vascular responsiveness and CBF to hypo- or hypercapnia will mean a greater change in brain (and central chemoreceptor) $\text{PCO}_2$ (and $[\text{H}^+]$) for any given change in arterial $\text{PCO}_2$, thereby increasing the...
slope of the \(\Delta V_{E}/\Delta P_{\text{aCO}_2}\) response and increasing controller gain above and below eupnea. The highly sensitive effects of changes in CBF on the control of eupneic ventilation, the ventilatory responsiveness to \(\text{CO}_2\), and the apneic threshold and \(\text{CO}_2\) reserve have been demonstrated experimentally during sleep in animal models using mechanical occlusion of carotid inflow (92, 93, 483), and in humans (735, 739) through the use of the cyclooxygenase inhibitor indomethacin to selectively depress cerebral vascular reactivity to \(\text{CO}_2\). Reductions in baseline CBF and in the cerebrovascular responsiveness to \(\text{CO}_2\) do occur with aging (50, 266), in severe OSA patients (133, 518), and with congestive heart failure (see sect. uG1). In turn, it is likely that these changes contribute to increases in controller gain and to the increased prevalence of sleep-induced ventilatory instability observed in these conditions. Limited clinical findings have shown that the treatment of congestive heart failure with the vasodilator captopril (an angiotensin-converting enzyme inhibitor) both increases CBF (537) and reduces apneic episodes (704).

G. Special Cases of Ventilatory Instability in Sleep

1. Chronic heart failure

The gradual waxing and waning of Cheyne-Stokes respiration (CSR) with cycling periods of 50–60 s duration occurs in about one-third of chronic heart failure (CHF) patients (460, 706) (see Fig. 1A). The causes of periodic breathing in CHF are multifactorial (281, 763). A key contributing abnormality is the increased controller gain in these patients, as defined by an increased ventilatory response slope to \(\text{CO}_2\), both above and below eupnea. The latter results in an absence of hyperventilation upon transition from wakefulness to sleep and a greatly reduced difference between eupneic \(P_{\text{aCO}_2}\) and the apneic threshold \(P_{\text{aCO}_2}\) during sleep (i.e., a narrowed \(\text{CO}_2\) reserve) (742). There are several potential sources of this increased controller gain as has been documented to occur in human patients and in animal models of CHF, including increased carotid chemoreceptor sensitivity (achieved in part via a reduced expression of nitric oxide and/or an increased expression of angiotensin II at the carotid chemoreceptor) (523, 656) and acute stimulation of lung vascular receptors via increases in left atrial and pulmonary vascular pressures (98, 633). In addition, a reduced cerebrovascular response to \(\text{CO}_2\) in CHF will 1) increase central chemoreceptor \(\text{CO}_2\) stimulation at any given level of raised arterial \(P_{\text{CO}_2}\), thereby enhancing the opportunity for ventilatory overshoot following apneas; and/or 2) reduce central chemoreceptor stimulation for any given reduction in arterial \(P_{\text{CO}_2}\), thereby enhancing the opportunity for apneas (738). A prolonged circulation time coincident with the reduced cardiac output in CHF will also delay corrective action by chemoreceptors. The resultant lengthening of apnea duration and increasing chemoreceptor stimulus levels enhances the opportunity for arousal, ventilatory overshoot and ventilatory periodicity (281).

So, even though chronic levels of hyperventilation and hypocapnia are common in CHF patients with CSR (280, 442), apparently (as with hypoxia) the stabilizing effects of the reduced plant gain attending the low \(P_{\text{aCO}_2}\) (see Fig. 7, top) are outweighed by the destabilizing effects of an increased controller gain (above and below eupnea) (see Fig. 7, bottom) and ventilatory instability prevails. Paradoxically, when eupneic \(P_{\text{aCO}_2}\) is driven even lower via a chronic ventilatory stimulus (acetazolamide), central sleep apnea and periodic breathing are significantly reduced in these patients (276). This is likely due to a further reduction in plant gain, with no change in controller gain, and therefore a widening of the protective \(\text{CO}_2\) reserve (436). Supplemental \(\text{O}_2\) also helps stabilize breathing in CHF probably by reducing controller gain (275, 279, 580). Predictably, even very small amounts of inhaled \(\text{CO}_2\) (or increased dead space) will prevent apneas and period breathing (299, 371), simply because the added \(\text{CO}_2\) will prevent the patients' arterial \(P_{\text{CO}_2}\) from falling below the threshold for apnea or hypopnea (53).

Finally, a new means of successfully preventing central apneas and periodicity in CHF uses a proportional assist servo-ventilator to provide a breathing with inspiratory pressure support (when a need is detected) together with a preset back-up respiratory rate to abort impending apneas (673).

Mixed apneas, i.e., central apneas followed by airway obstruction within the same apneic event, are common in CHF for significant portions of sleep time. This is particularly true in those patients who are obese and/or with a history of snoring, suggesting a high likelihood of a high passive airway \(P_{\text{crit}}\) (278, 282, 682). Accordingly, CPAP treatment also reduces apnea and CSR in some but not all CHF patients (277), possibly because 1) airway narrowing and obstruction are prevented, thereby removing one potential cause of blood gas changes, chemoreceptor stimulation, and arousal that will elicit ventilatory overshots (262); 2) preventing the reflex central apneas caused by airway closure during early expiration (224, 576); and 3) reducing ventricular afterload and pulmonary vascular pressure.

2. Hypoxic-induced periodic breathing

Sojourners to high altitude commonly experience restlessness and nonrefreshing sleep, in part due to the common occurrence of periodic breathing. During NREM or REM sleep hyperventilation begins immediately upon hypoxic exposure and intensifies with time (see Fig. 1B) (53, 54). After ~10 min of hypoxia in the sleeping human,
tidal volume begins to oscillate in a waxing and waning pattern. These oscillations keep increasing in magnitude as hypoxia is maintained and PaCO₂ falls further to the level of the apneic threshold. Commonly then an augmented inspiration occurs and the subject begins overt periodic breathing cycles of ~15–25 s duration, characterized by two or three huge tidal volumes followed by apneas of 5–15 s duration as well as large swings in cerebral blood flow (130). During these periodic cycles arterial SaO₂ swings wildly along the steep part of the oxygen dissociation curve.

As with CHF, the principle reason for apnea and periodic breathing in hypoxic sleep is likely to be the increased controller gain, as evidenced by the steep increase in CO₂ response slope above and below eupnea and the greatly narrowed CO₂ reserve (436, 742). Accordingly, during apnea elicited via very minimal amounts of transient hypcapnia (~1–2 mmHg less than eupnea), PaCO₂ rises commensurate with sharp reductions in PaO₂ below an already hypoxemic baseline, and the interaction of these asphyxic stimuli greatly enhances carotid chemoreceptor activity and the drive to breathe. Upon rapid restoration of normoxic SaO₂ via increased FiO₂, periodic breathing continues with prolonged apneic periods until hyperventilation is gradually reduced and PaCO₂ returns to normal.

Unlike the gradual waxing and waning of CSR in CHF, periodic breathing in hypoxia occurs in breath “clusters” with tidal volume increasing from zero to three to four times control, almost instantaneously following each apnea. We believe this implies the presence of a transient arousal at apnea termination which would further augment the responsiveness of the respiratory control system and produce the sudden ventilatory overshoot (177, 301). Although cortical EEG arousals have not been consistently observed throughout periodic breathing in hypoxia (53, 301), it is still feasible that arousal only at the level of the brain stem (249) could greatly magnify this chemoreceptor responsiveness. An additional as yet untested influence in this dynamic, physiologically complex condition would be brain hypoxia acting independently as a ventilatory stimulant (112, 154, 625), an effect which may be sensitized via a simultaneously enhanced carotid chemoreceptor input (see sect. mF2).

The amount of periodic breathing in sleep is greatly reduced over time in hypoxia (53, 54) and when chronic respiratory stimulants (such as acetazolamide or doxapram) are administered to sleeping sojourners at high altitude (547, 659). Perhaps then, in short-term hypoxia, the stabilizing effect of a reduction in plant gain associated with a reduced PaCO₂ is offset by the marked effect of hypoxia on increasing controller gain and therefore loop gain, whereas with acclimatization or superimposed chronic stimuli, the further reductions in PaCO₂, and plant gain override the increased controller gain. Additionally, a few weeks of intermittent hypoxic exposure in canines produced no aftereffects on ventilation 1 day following the cessation of exposure, but did decrease the apneic threshold and widen the CO₂ reserve in the sleeping animal (294).

3. Opioid-induced periodicity

Periodic, cluster-type, ataxic breathing patterns have been reported with high prevalence during NREM sleep in patients administered chronic doses of opioid medications (703, 705). The severity of apneic events is dependent on opioid dose (703), occurs predominantly in NREM rather than REM sleep (278) and is only very rarely associated with chronic (daytime) CO₂ retention (672). So, while acute opioid administration in animals and awake humans is well known to cause hyperventilation and apnea via mechanisms acting at the level of the medullary respiratory rhythm generator (331, 614), with chronic opioid administration sleep appears to be required to unmask the instabilities in ventilatory control (also see sect. mF). Perhaps the central depressant effects of opioids might sensitize the apneic threshold and narrow the CO₂ reserve below eupnea. As with other cases of background hypventilation, plant gain would be enhanced and the CO₂ reserve narrowed. However, as with hypoxia, there remains no ready explanation for the abrupt transitions from apnea to transient hyperventilation. Like other forms of predominately central sleep apneas, opioid-induced periodicity is treatable via servoventilation (278). Major clinical concerns over opioid-induced sleep apnea are twofold: 1) the dramatic increase in the therapeutic use of opioids (such as methadone, oxycodin, and morphine) over the past decade (519) and 2) the high mortality rates reported for these patients, especially the many sudden deaths occurring in the early morning or in bed (278).

In summary, the causes of cyclical periodic breathing containing predominantly central or mixed apneas are multifactorial, and while we can usually point to specific contributing factors such as those affecting plant and/or controller gains, it is not yet possible to predict with certainty exactly how deficiencies in different elements contained within the control system, especially chemoreceptor control, mesh to produce this repetitive series of ventilatory under- and overshoots in the sleeping state. Since breathing in the sleeping state is so very critically dependent on chemoreceptor control, it is imperative that we follow recent leads from evidence in reduced preparations to more fully understand how peripheral and central chemoreceptors influence each other’s responsiveness and in turn overall ventilatory responsiveness in the integrated, intact preparation studied across varying states of consciousness.
H. Interaction of Neurochemical Control Mechanisms and Upper Airway Anatomy in OSA

We now attempt to integrate our previous discussions of airway anatomy, neurochemical control of upper airway dilators, chest wall pump muscles, and ventilatory and sleep state stability into a more cohesive understanding of the pathogenesis of cyclical, repeated airway obstructions. First, there is little doubt that compromised upper airway anatomy plus the loss of wakefulness input increases passive $P_{\text{crit}}$, thereby rendering the airway highly susceptible to closure at sleep onset. However, the question remains as to why obstructive sleep apneas recur in a cyclical pattern of ventilatory undershoots and overshoots in OSA patients. Several observations have implicated deficits in neurochemical control and stability of central respiratory motor output and upper airway neuromuscular recruitment as key contributors to cyclical OSA. For example, fluctuations in chemical stimuli, respiratory drive, and ventilation are associated with reciprocal oscillations in EMG and airway resistance, with the airway narrowing the most when drive is at its nadir (8, 23–25, 256, 464, 708). Accordingly, very early in the course of a hypocapnic-induced central apnea, bronchoscopic imaging studies in sleeping humans show airway narrowing and often complete closure in the absence of changes in sleep state or in inspiratory effort (25, 324) (see Fig. 5).

Many patients with severe OSA show a significantly higher than normal loop gain for ventilatory control, as determined by the propensity for periodic breathing observed during ventilatory assist (714, 752) or a greater ventilatory response to single breaths of CO$_2$ (257). Loop gain correlates with OSA severity best in those patients with a passive $P_{\text{crit}}$ that is near atmospheric pressure, i.e., not too negative or too positive (714) (see also scenario 2 below).

A significant portion of subjects with positive passive $P_{\text{crit}}$ (i.e., highly collapsible airways) have a relatively low AHI, and many of these patients with high mechanical loads on their airway maintain airway patency for significant periods of time throughout sleep without experiencing repeated, cyclical obstructions (714, 752, 755).

Normal subjects decrease their $P_{\text{crit}}$ during sustained reductions in airway pressure (i.e., below $P_{\text{crit}}$ levels determined under passive conditions), whereas many OSA patients do not, implying that the threshold for upper airway muscle activation in response to increased chemostimulation is much higher in the OSA patient (492, 493).

Many OSA patients undergoing either tracheostomy (465) or CPAP treatment (193, 676) continue to show periodic ventilatory cycling for variable periods of time. Although often difficult to detect via routine, noninvasive polysomnography, “mixed”, i.e., central followed by obstructed apnea within the same event (see Fig. 1C), are common, and the same subject will often experience central and predominantly obstructed apneas within the same night (682).

Animal and human studies show a linear chemorecceptor-driven recruitment of diaphragm EMG as opposed to a highly alinear, threshold-like response of upper airway muscle EMG to increasing chemoreceptor stimuli (227, 251, 370, 517).

In summary, these observations point to strong links between neurocontrol mechanisms and airway obstruction, but the determinants of these links are multifactorial. Accumulating evidence obtained in sleeping patients with highly variable magnitudes of airway mechanical loads points to two scenarios that illustrate the potential importance of compensatory neuromuscular control mechanisms on the one hand and central control instability on the other in determining cyclical OSA, when either or both occur in persons with upper airways that are anatomically susceptible to narrowing and closure. For clarity, we present these examples separately, but clearly the underlying causative mechanisms are common to both scenarios of cycling OSA.

1. Scenario 1: compensatory responses to airway obstruction determine susceptibility to OSA cycling

This scenario is illustrated in Figure 8 from Younes (753) in which CPAP pressure in a sleeping OSA patient is suddenly reduced to less than $P_{\text{crit}}$ (passive), thereby creating an obstructive apnea. This experimentally imposed obstruction sets the stage to consider the influence of compensatory neuromuscular mechanisms in coping with the resolution of an airway obstruction in response to rising chemoreceptor sensory input. As previously discussed and also recently reviewed (754), these mechanisms include 1) ability to effectively recruit upper airway dilators during the apnea and prior to arousal; 2) effectiveness in converting the neural drive into dilator muscle shortening and airway reopening; 3) arousal threshold, which will vary with a) sleep state, b) the magnitude of sensory input from chemoreceptors, and c) the patient’s sensitivity to a given chemosensitive input to the higher CNS; 4) the magnitude of the ventilatory overshoot once airway patency is reestablished, as determined by the rate of rise of chemoreceptor stimulus magnitude and response sensitivity (i.e., controller gain). Transient arousals are major determinants of controller gain and therefore the magnitude of the ventilatory overshoot at end-apnea; and 5) the subsequent ventilatory undershoot will determine the magnitude of the reduction in respiratory motor output to both airway and pump muscles in response to the inhibitory influences of the hypocapnia resulting from the preceding ventilatory overshoot. These factors will vary in magnitude and relative importance among subjects and even within the same subject throughout the night. It is
proposed that the nature of the interaction among these factors within and immediately following each obstructive event will determine whether initial obstructive events are followed by stable breathing, slow evolving hypopneas with occasional arousals, or repetitive obstructions (754).

2. Scenario 2: control system instability determines OSA cycling

A second scenario coupling neuromechanical control mechanisms to cyclical OSA is shown in Figure 9 from Warner et al. (708), which illustrates how oscillations in respiratory motor output (as imposed experimentally in this study by brief hypoxic exposures) may lead to cyclical airway obstruction. A subject is shown in whom airway resistance was already substantially elevated during sleep, but the airway was still patent, prior to imposition of the hypoxia. During the hypoxic-induced ventilatory oscillation period, airway caliber narrowed and then closed at the nadir of central respiratory motor output. This snoring subject likely had a passive pharyngeal closing pressure very close to atmospheric, which is characteristic of groups of subjects who show high, positive correlations between loop gain and AHI (714). In other subjects whose sleep-induced increase in airway resistance was barely measurable (i.e., highly negative passive $P_{crit}$), imposing oscillations in respiratory motor output had no or very little effect on airway resistance, and in patients with highly positive passive $P_{crit}$, cyclical airway obstruction occurred during sleep without the need to superimpose further ventilatory instability.

A similar pattern of cyclical obstruction related to an oscillating respiratory drive is prevalent in anatomically susceptible elderly subjects (258, 477). However, in the elderly an unstable sleep state, especially in light stages I and II, is prevalent (477) and appears to be the primary driver to ventilatory periodicity. In these subjects, changing chemoreceptor stimuli would likely also eventually contribute to the periodicity in airway caliber, but only in a secondary role.

3. Summary: treatment implications

All of the causes of these couplings of neural control mechanisms to the severity of cyclical OSA have not yet been determined, although there have been many attempts especially recently to study increasing numbers of OSA patients during sleep (summarized in Refs. 714, 715, 754). The propensity for cyclical OSA will vary within a subject even within the same night as, for example, 1) the majority of all OSA patients will have significant periods during the night of stable breathing without arousals or ventilatory overshoots or undershoots, indicating that they have compensated quite effectively for increases in mechanical loads on the airway; or 2) unstable ventilatory control may result in complete airway obstruction (apnea) or only partial airway narrowing (hypopnea), depending on the magnitude of the oscillating chemical drive, the chemosensitivity to inhibition and excitation of both upper airway and chest wall pump muscles in response to oscillations in chemoreceptor stimuli and the patient’s inherent passive $P_{crit}$.

So several interrelated mechanisms clearly contribute to a patient’s success or failure in smoothly exiting an obstructive apnea to resume stable breathing or to their experiencing cycling obstructive behavior; or to their propensity to experience oscillations in central respiratory...
motor output which result in repeated airway obstructions. Accordingly, at this point in our understanding, it seems clear that 1) cycling OSA requires anatomical predisposition to airway closure because no amount of central respiratory motor output instability will result in closure or substantial airway narrowing in normal subjects with highly negative passive $P_{\text{crit}}$; and 2) an airway susceptible to closure does not guarantee cycling OSA, because compensatory neuromechanical control mechanisms are an equally important determinant of this repetitive OSA cycling.

Recognizing these complex anatomical-functional relationships in general and recognizing how $P_{\text{crit}}$, dilator muscle recruitment, ventilatory response gains, and arousability vary among individual patients may provide clues to OSA treatment by means other than CPAP or by other physical means of manipulating airway caliber, per se. Certainly CPAP clearly prevents airway obstruction in OSA patients; however, it is not tolerated well or at all by some, compliance is low in many others, a residual periodic breathing may ensue in many CPAP users and in some CHF patients CPAP may be contra-indicated (66, 118, 277, 317). To date, attempts to treat OSA with such single entities as increased $F_{\text{ICO}_2}$ or $F_{\text{IO}_2}$, pharmacological respiratory stimulii, or sedatives (to diminish arousability) have met with very limited success (234, 257, 754). Future attempts in this regard should focus on targeting specific treatments to patients with specific deficits in control system stability or compensatory capabilities in the recruitment of upper airway muscle dilators. Some limited findings provide a basis for future studies, especially in OSA patients with only moderately elevated passive $P_{\text{crit}}$ (708, 714). For example, use of supplemental $F_{\text{IO}_2}$, which is known to reduce $CO_2$ sensitivity above and below eupnea and to increase the $CO_2$ reserve below eupnea (740), was shown to reduce AHI substantially in some OSA patients with high loop gain (715). Furthermore, increasing respiratory motor output via increased $F_{\text{ICO}_2}$ stabilizes breathing and greatly reduces upper airway resistance in sleeping subjects with partially obstructed airways (23, 24). Alternatively, pharmacologically induced stimulation of respiratory motor output via such agents as carbonic anhydrase inhibitors (276) or 5-hydroxytryptamine (5-HT)$_{1A}$ receptor agonists (724, 745) reduce plant

![FIG. 9. Determinants of how imposed oscillations in respiratory motor output and ventilation (via hypoxia) might lead to cyclical airway obstruction in a subject with an upper airway anatomy susceptible to closure during sleep. Mean values are shown for upper airway resistance during wakefulness and NREM sleep at the left. Breath by breath peak upper airway resistance ($R_{aw}$) is then shown before, during, and after hypoxic exposure. A subject with a fivefold increase in $R_{aw}$ from awake to sleep but with stable breathing and mild $CO_2$ retention is shown. During early hypoxic exposure, oscillations in respiratory muscle output (EMG$_{di}$) but without central apnea occurred, leading to periodic airway obstruction coincident with the nadir of EMG$_{di}$. However, with continued hypoxia and fully developed periodic breathing with apneas and profound $O_2$ desaturation and $CO_2$ accumulation, $R_{aw}$ remained very low (even approximating waking levels) during breaths with high levels of respiratory motor output following each central apnea. Then, upon return to normoxia, cyclical airway obstructions returned again at the nadir of EMG$_{di}$. [From Warner et al. (708).]]
gain and ventilatory instability. Perhaps application of these agents to patients with only moderately elevated passive $P_{\text{crit}}$, combined with high loop gain, may well reduce cycling airway obstructions (708, 714).

IV. NEUROCHEMICAL CONTROL OF UPPER AIRWAY PATENCY

A. Overview

Collectively the very characteristics rendering the pathophysiology of obstructive sleep apnea support the concept that this disorder should be readily amenable to pharmacotherapeutics. First and foremost, obstructive sleep apnea is a remarkably focal process. The site of primary collapse typically lies within a small segment of the oropharynx, where collapse is a consequence of sleep state-dependent reductions in tone in specific upper airway dilator muscles. Second, individuals with obstructive sleep apnea have the capacity in wakefulness to stent open the upper airway through proper activation of the upper airway dilators; it is only in sleep, when neurochemical drive to upper airway motoneurons is altered, that insufficient muscle activity ensues, resulting in pharyngeal collapse. Thus readily reversible changes in neural drive to the upper airway dilator muscles underlie the pathogenesis of obstructive sleep apnea (also see sect. 11.E). Changes in neural drive imply alterations in the amounts or types of neurotransmitters delivered to upper airway motor nuclei. Together, these insights into the pathophysiology of obstructive sleep apnea highlight the importance of sleep state-dependent neurochemical changes reducing the activity of upper airway dilator motoneurons. Over the past two decades tremendous progress has been made in identifying the relevant muscles and their motoneurons and in elucidating the neurochemical control of the upper airway.

B. Upper Airway Dilator Motoneuronal Groups

Multiple muscle groups are involved in the maintenance of upper airway patency in persons anatomically predisposed to obstructive sleep-disordered breathing. Because the oropharynx is so highly collapsible from multiple directions, most individuals with a predisposition to sleep-related collapse of the upper airway rely on opposing muscle groups to work in unison to prevent upper airway collapse, as summarized in Figure 10. Thus it is important to understand the neurochemical control of multiple dilator muscle groups and how the muscles work together. Unfortunately, to date, a tremendous emphasis has been placed on understanding neurochemical control of the genioglossus, or tongue muscle, and its major

![Fig. 10.](image-url)
nerve, the hypoglossal. The genioglossus is the largest and most powerful upper airway dilator. However, genioglossus muscle activation alone may be insufficient to reduce pharyngeal collapsibility (386, 461, 692). Therefore, many of the neurochemical studies of hypoglossal neurochemical control presented below will need to be repeated for the other important dilator groups with different vectors. In summary, then, in individuals with collapsible upper airways, important dilating forces are contributed by several of these groups of motoneurons: motor trigeminal (V), facial (VII), glossopharyngeal (IX) motor vagus (X), and hypoglossal (XII). Muscles regulating lung volume and neck and jaw position will also contribute to upper airway patency (236), and thus cervical ventral horn motoneurons may also influence upper airway patency.

C. Multiplicity of Function for Upper Airway Motoneurons

One of the greatest challenges in designing therapies to activate upper airway motoneurons is that these muscles also serve critical functions for speech, mastication, propulsion of food into the esophagus, and airway clearance. Thus any attempt to grossly activate one or more upper airway muscles tonically is likely to substantially interfere with these other vital pharyngeal tasks. Central pattern generators are shared for swallowing, sneezing, and breathing (399). Additionally, pharyngeal muscles must also coordinate with ventilatory pump muscle activation. Specifically, the upper airway should be stented open prior to the development of negative intraluminal pressures within the pharynx from activation of the diaphragm and other pump muscles. The coordination of all of these complex processes ultimately merges at pharyngeal motoneurons. Thus an understanding of the neurochemical mechanisms by which sleep state alters resting resultant negative intraluminal pressures, utilizes rapid onset/offset with an excitatory neurotransmitter, glutamate, and inhibitory neurotransmitters, GABA and glycine. Receptors for the primary excitatory neurotransmitter are \( \alpha \)-amino-3-hydroxy-5-methyl-4-isoxazole propionic acid (AMPA), kainite (KA), and \( \alpha \)-methyl-\( \alpha \)-aspartate (NMDA). Specific receptor subtypes for each of the glutamate receptor groups have been identified for brain stem motoneurons (392, 504). Yet, the relative roles for each receptor subtype in excitation of motoneurons and premotoneurons is just beginning to be explored. Several of these receptor subtypes can be rapidly desensitized by intense activation and/or redox alterations (365). Clearly, this is an important area that deserves further study and one that may contribute to progression of apneic events across the night. In addition to fast action of glutamate at ionotropic receptors, glutamate serves as an excitatory neuromodulator by way of an array of metabotropic receptors.

GABA and glycine are the primary inhibitory neurotransmitters acting at motoneurons, and functional GABA\(_A\) and glycine receptors are evident on most upper airway dilator motoneurons (480). Glycine plays an essential role in REM sleep postural atonia (631, 733), but the roles of GABA and glycine in atonia of hypoglossal motoneurons are controversial (746). Morrison and co-workers (431, 432) tested the relative contribution of GABA and glycine to genioglossal muscle suppression in an adult rat model with spontaneous sleep and a chronic dialysis probe in the hypoglossal nucleus. Antagonists to both inhibitory neurotransmitters increased baseline nerve activity but did not show a preferential increase for REM sleep, suggesting that neither neurotransmitter contributes significantly to genioglossal muscle atonia in spontaneous REM sleep (368, 431, 432). Of clinical interest, there is a case study of strychnine (a glycine antagonist) effects on genioglossus and tensor veli palatine muscle activity in an individual with obstructive sleep apnea. In this case, strychnine markedly increased tensor veli palatine activity and also increased genioglossus activity relative to arterial oxygen tension (548). For an hour of sleep (at
maximum drug effect), apneas were completely abolished and ventilatory efforts were remarkably regular. Thus glycine may not have significant effects on sleep-related respiratory suppression in animals without collapsible airways, but in humans with obstructive sleep apnea, a role of glycinergic tone in sleep-related suppression of muscle tone may be a pharmacological tool for sleep apnea and deserves further study. It is conceivable that glycinergic receptors on upper airway motoneurons could be altered with gene therapy into pharyngeal muscles without disturbing other pharyngeal functions.

2. Neuromodulators

A) SEROTONIN EFFECTS AT UPPER AIRWAY MOTONEURONS. Serotonin is a powerful modulator of motoneuronal activity (308, 721) and is, perhaps, the best studied for its role in upper airway motoneuronal activity. Activity of the neurons that deliver this neuromodulator throughout the brain have highest firing rates in wakefulness; rates are reduced in NREM sleep, and these neurons are relatively quiescent in REM sleep (243, 696). Leszek Kubin hypothesized that reductions in upper airway dilator muscle activity might be the result of sleep state-dependent decline in serotonin delivery to dilator motoneurons (320). In the decerebrate, vagotomized cat, there is endogenous serotoninergic tone in hypoglossal motoneurons (321). Richard Horner confirmed significant intrinsic serotoninergic activity in the hypoglossal nucleus in the anesthetized, vagotomized rat (636) yet found little intrinsic serotoninergic excitation in the intact, awake rat (637). The Horner group also showed that vagotomy increases serotoninergic tone in the hypoglossal nucleus and genioglossus activity; in contrast, in an adult rat with intact vagi, there is little serotoninergic tone. (637). Thus animals with unobstructed breathing in sleep may have little to no endogenous 5-HT contributing to resting upper airway motoneuronal tone. Nonetheless, a consistent finding across
research labs is the ability of exogenous 5-HT to augment upper airway nerve or muscle activity.

Serotonergic neuromodulation of upper airway motoneurons in individuals with obstructive sleep apnea, however, may differ from the role serotonin plays in animals without obstructive sleep-disordered breathing. Specifically, upper airway muscle activity may be recruited in wakefulness to stent open the upper airway in individuals with obstructive sleep apnea (405). An established animal model with spontaneously occurring sleep state-dependent collapse of the upper airway is the English bulldog (238). In NREM sleep, the dog has hypopneas (partial obstructions, with oxyhemoglobin desaturations and arousals); in REM sleep the dog has significant apneas with more pronounced reductions in arterial oxyhemoglobin saturation (238). Veasey et al. (696) examined the importance of serotonin in the maintenance of the upper airway in the English bulldog, by examining rapid sequencing computerized tomography of the upper airway before and after systemic administration of two serotonin antagonists, methysergide and ritanserin (696). Administration of the serotonin antagonists resulted in temporary collapse of the upper airway (measured with cinematic computerized tomography), associated with reduced upper airway muscle activity and oxyhemoglobin desaturations, without electrographic or behavioral evidence of sleep. Thus, in an animal model of obstructive sleep-disordered breathing, serotonin plays an important role in maintenance of pharyngeal patency. It is important to acknowledge that because the drugs were administered systemically, it is difficult to discern the site of action for the serotonin antagonists. Nonetheless, administration of broad-spectrum serotonergic agonists largely prevents airway collapse in the English bulldog in NREM sleep, supporting the concept that serotonergic neuromodulation could be used to treat obstructive sleep apnea. In REM sleep, serotonergic agents reduced sleep-disordered breathing events in the bulldogs, but are less effective, particularly in preventing events in phasic REM sleep (695).

B) CLINICAL TRIALS OF SEROTONERGIC AGENTS IN OBSTRUCTIVE SLEEP APNEA. Both serotonergic and serotonin receptor antagonist drugs have been tested for effectiveness in reducing obstructive sleep-disordered breathing, including 5-methyltryptophan, fluoxetine, paroxetine, trazodone, and mirtazapine. In summary, none of the drugs tested has proven universally effective (52, 222, 313, 573, 590). In addition to the excitatory hypoglossal motoneuronal postsynaptic effects of 5-HT2A R and possibly 5-HT2C R, there is an important inhibitory effect observed with 5-HT1B R receptor activation (459). This effect is blockade of glutamate release (62, 618, 619). The effectiveness of 5-HT1B antagonists in obstructive sleep apnea has not been tested, despite the above studies supporting 5-HT1B R as a potential pharmacological target.

C) VENTILATORY EFFECTS OF SEROTONERGIC AGENTS BEYOND UPPER AIRWAY MOTONEURONS. In considering serotonergic therapies for obstructive sleep apnea, it is important to consider where else activation of various 5-HT receptor subtypes might impact on ventilatory drive to upper airway muscles and overall ventilation (for review, see Ref. 48). 5-HT also has direct effects on medullary premotoneurons. Activation of the 5-HT2 R receptor subtypes with iontophoretic application of α-methyl-5-HT depolarizes medullary expiratory and postinspiratory neurons, resulting in a reduction of the persistent and postsynaptically activated potassium currents (333). Systemic administration of a partially selective 5-HT1A antagonist, 8-OH DPAT, increases respiratory drive, while ventilation can be suppressed by 5-HT1A antagonist drugs (671). This effect was recently shown to be centrally mediated (662). Like 5-HT2A R, 5-HT1A activation promotes wakefulness. Thus 5-HT1A agonists may be helpful with anesthesia and analgesia suppression of ventilation but would likely disrupt or reduce sleep if administered for sleep apnea. The pre-Bötzinger region contains critical respiratory pattern generator neurons in mammals, and at this site, 5-HT can influence ventilatory drive. Within this respiratory rhythm generator, 5-HT1AR is the predominant 5-HTR. 5-HT1AR agonists applied directly to these respiratory rhythm neurons suppress apneusis and cause a pattern of markedly prolonged inspiratory efforts, and this has been translated into clinical medicine (332, 724). This breathing pattern may occur in barbiturate overdoses, and 5-HT1AR agonists, including buspirone, a partial 5-HT1AR agonist, can reverse the drug-induced apneusis. This partial agonist does not, however, reverse narcotic-induced respiratory suppression in humans (458). Pharmacological blockade of the 5-HTR within the pre-Bötzinger region also directly affects ventilation (554). A highly selective 5-HT2R antagonist, CB113808, induces central apneas when injected into the pre-Bötzinger area, while a selective agonist, BIMU8, increases respiratory drive and importantly can reverse narcotic-induced apnea without reversing the analgesic effect of narcotics (554). Thus 5-HT2R agonists may have utility in managing opioid-induced ventilatory suppression, but could also have potential for sedative- and sleep-induced ventilatory suppression. The third 5-HT subtype with activity in the pre-Bötzinger region that influences abnormal breathing is the 5-HT2A R that is critical for postischemic gasping and recovery of suppressed ventilatory drive. It is imperative as we explore potential pharmacological agents that we take into consideration all possible sites of action and expected effects. For example, 5-HT2A R activation can worsen asthma, hypertension, psychoses, and thromboembolic disease. Thus, while 5-HT2A R activation can increase upper airway dilator tone, 5-HT2A R agonists are not likely to be safe and well-tolerated (352).
5-HT\textsubscript{1A} effects on ventilation are present in several additional sites within the central nervous system. In the hypothalamus, there are 5-HT\textsubscript{1A} effects on the ventilatory response to hypoxia. Here, activation of 5-HT\textsubscript{1A}R inhibits the increased ventilation response to hypoxia. This effect is also observed with active 5-HT\textsubscript{3} compounds (186). Activation of the 5-HTR in the nucleus tractus solitarius in the rat increases ventilatory frequency (539). In addition to 5-HT\textsubscript{1A} activation of ventral respiratory neurons, described above, there is also an excitatory effect of 5-HT\textsubscript{1A}R activation in the dorsal motor vagal nucleus (76). Specifically, injection of the 5-HT\textsubscript{1A} agonist 8-OH DPAT into the dorsal motor nucleus increases ventilation without altering upper airway motoneuronal activity.

Serotonergic agents have significant ventilatory effects within the peripheral nervous system. 5-HT applied to the nodose ganglion suppresses respiration and induces apneas (751). 5-HT\textsubscript{3} is also responsible for suppression of ventilatory drive at the nodose ganglion (751). Administration of a 5-HT\textsubscript{3} antagonist, ondansetron, to normal rats reduced the number of central sleep apneic events (82), and this effect is clearly a peripheral effect, consistent with activation of the nodose ganglion (327). Oral administration of this same 5-HT\textsubscript{3} antagonist to the English bulldog reduced the number of sleep-disordered breathing events in REM sleep, without affecting NREM sleep events (693). Therefore, 5-HT\textsubscript{3} antagonists may be effective in reducing REM sleep events in some individuals with OSA with sleepiness (OSAS). A recent clinical trial in individuals with OSAS did not observe an overall effect of ondansetron (single dose) on OSAS events (651).

D) SUMMARY. Serotonin delivered to the upper airway motor nuclei in all animal models tested can increase hypoglossal nerve/genioglossus muscle activity across sleep states. Whether serotonin can augment the activity of all of the motoneurons necessary to stent open the upper airway requires further investigation. Although it is disappointing that the excitatory 5-HT receptor subtypes on upper airway motoneurons are not suitable targets for systemically administered therapies, there may be several ways to locally enhance 5-HT within motoneurons to a level sufficient to maintain critical muscle activity in sleep. In all likelihood, there will be distinct subsets of patients with obstructive sleep apnea who will respond differentially to serotonergic agents, and these groups need careful phenotyping in future drug studies testing effectiveness in sleep apnea.

E) NOREPINEPHRINE. Norepinephrine has several similarities with 5-HT as a neuromodulator of motoneuronal function and also has several important distinctions. Like 5-HT, norepinephrine has an overall excitatory effect on hypoglossal nerve function when delivered into the hypoglossal nucleus (5, 89, 182, 486), and like 5-HT, noradrenergic neurons show reduced firing rates in NREM sleep and are relatively quiescent in REM sleep (18). Hypoglossal motoneurons are innervated by the A7 noradrenergic group, which shows a similar state dependency in firing (20). In contrast to minimal endogenous 5-HT effects in the hypoglossal nucleus in spontaneously sleeping animals, endogenous norepinephrine contributes to genioglossus tone in wakefulness and in NREM sleep (89). Microinjection of a noradrenergic antagonist into the hypoglossal nucleus reduces genioglossus muscle activity by 25–50%, supporting intrinsic noradrenergic tone in the hypoglossal motor nucleus in both wake and NREM sleep. In contrast, in REM sleep, the noradrenergic antagonist has no effect, suggesting that in REM sleep, there is little to no noradrenergic tone in the hypoglossal nucleus. Moreover, norepinephrine contributes to several important upper airway reflexes (138), while 5-HT does not appear to contribute measurably. Specifically, systemic administration of prazosin, an adrenoreceptor \( \alpha_{\text{1B}} \) antagonist, prevents the masseter muscle activation to multiple stimuli (termed the jaw closure reflex), while a broad-spectrum serotonin antagonist does not alter this reflex response (642). \( \beta \)-Adrenergic tone contributes to motoneuronal responses to either chemoreflexes or trigeminal diving and superior laryngeal nerve reflexes (209). Whether noradrenergic tone in upper airway dilator motor nuclei contributes to the negative pressure airway reflex should now be examined, and this examination should include, not only hypoglossal, but motor trigeminal and facial nerve responses. The only adrenergic receptor subtype mRNA expressed in the majority of hypoglossal motoneurons was for the \( \alpha_{\text{1B}} \) receptor (701). The functional significance of the \( \alpha_{\text{1B}} \) receptor has been established as a powerful excitatory effect present under basal conditions for both hypoglossal and trigeminal motoneurons (89, 162, 164, 642). Thus \( \alpha_{\text{1B}} \) agonists may have a role in select patients with mild NREM sleep predominant sleep apnea. In light of the excitatory effect at the \( \alpha_{\text{1N}} \) receptor, there is a concern that prazosin could worsen sleep apnea.

Few clinical trials have examined the effects of noradrenergic agents on obstructive sleep apnea. Clonidine, an \( \alpha_2 \) agonist antihypertensive, suppresses the activity of noradrenergic neurons, yet the drug also suppresses REM sleep. In light of the REM sleep suppressant effect, clonidine was examined as a potential treatment for sleep apnea in one placebo-controlled trial (\( n = 8 \)) where two placebo polysomnographies were compared with two clonidine polysomnographies for each subject (273). Overall, 0.2 mg clonidine at bedtime suppressed REM sleep by 40%, had no effect on NREM sleep apnea hypopnea events, but by suppressing REM sleep reduced the total number of REM sleep events. Whether clonidine has efficacy in patients with predominantly REM sleep apnea deserves further study. Two noradrenergic reuptake inhibitors have been tested for effects on sleep apneic events, protriptyline and a newer agent atomoxetine (37,
In the earlier study, protriptyline at 10 mg/day was administered for 4 wk and then examined for its effects on apnea frequency. The apnea index dropped for the group from 57 ± 9 to 33 ± 8, \( P < 0.05 \). The improvement was only present in NREM sleep, but REM sleep was markedly suppressed (222). In the more recent trial, atomoxetine was administered as 40–80 mg for 4 wk prior to polysomnography in 12 subjects with mild sleep apnea (5–15 apneas-hypopneas/h). Although subjective sleepiness improved, there was no improvement in the apnea/hypopnea index (37).

F) Adenosine and ATP. Obstructive sleep apneic events (through hypoxia) are expected to alter motoneuronal and extracellular levels of purines: reducing ATP and increasing adenosine. Adenosine modulates motoneuronal activity, where the overall effect of adenosine injected into the hypoglossal nucleus suppresses hypoglossal motoneuronal activity (516). This hyperpolarization effect is mediated through the \( A_1 \) adenosine receptor and is thought to reduce glutamatergic signaling, as evidenced by \( A_1 \) agonist-induced reduction in the amplitude of excitatory postsynaptic potentials (46). Like adenosine, ATP modulates glutamatergic neurotransmission (356). The pharmacology of ATP neuromodulation involves >18 ion-gated and \( G \) protein-coupled receptor subtypes, and only a few of these have been examined for upper airway dilator motoneurons. Funk et al. (181) examined the neuromodulatory effects of ATP on hypoglossal motoneuronal activity in both medullary slices and in anesthetized adult rats, observing an excitatory effect in both models. This excitatory effect was mediated at least in part by the \( P_2 X_2 \) ATP receptor subtype on hypoglossal motoneurons. An excitatory modulation has also been observed for phrenic motoneurons; however, this effect is more complex and is followed by a secondary inhibition of phrenic motoneurons, suggesting the involvement of other purinergic receptors (406). Clearly, the pharmacology of the secondary inhibitory effects should be further developed.

G) Acetylcholine. Acetylcholine (ACh) injected directly into the hypoglossal nucleus markedly suppresses hypoglossal nerve activity in medullary slice preparations (88) and genioglossus muscle activity in anesthetized adult rats (367). While the overall effect is inhibitory, activation of select pontomedullary cholinergic neurons can activate or suppress hypoglossal motoneurons. A very thorough description of these opposing effects was recently described (319). Intrinsic ACh inhibitory activity is supported by the observation that administration of an acetylcholinesterase inhibitor into the hypoglossal nucleus suppresses activity (367). Pontine and forebrain cholinergic neurons are least active in NREM sleep, most active in REM sleep, and have intermediate activity in wakefulness (698). The sleep state dependency of medullary cholinergic neurons innervating brain stem motoneurons (563, 684) has not been described. ACh targets two distinct groups of receptors: muscarinic and nicotinic receptors. The inhibitory effect of ACh on hypoglossal motoneurons is muscarinic and can be largely blocked with a muscarinic antagonist, atropine (367). The muscarinic inhibitory effect is likely presynaptic suppression of glutamate release (47). This muscarinic suppression of hypoglossal activity may be one important mechanism by which morphine suppresses genioglossus muscle activity, as dialysis of morphine into the hypoglossal nucleus increases ACh release, and this local increase results in suppression of hypoglossal nerve activity (622). However, there is also an excitatory effect of ACh through activation of nicotinic receptors that is masked by the overwhelming muscarinic effect (367). The nicotinic receptor subtypes include the \( \alpha 4\beta 2 \) and \( \alpha 7 \) receptor subtypes (536). Of clinical significance, there is a rapid desensitization of nicotinic receptors in the hypoglossal nucleus within minutes (88, 536).

The first clinical trial of an atropine-like substance involved testing the effectiveness of belladonna on sleep-disordered breathing in infants 4–46 wk of age (289). In this report, apneas were prevented fully in 10 of 15 infants. To date, this has never been reexamined. There are several studies testing the effectiveness of nicotine on obstructive events in adults. In the two controlled trials, transdermal nicotine disrupted sleep without improving the frequency of obstructive breathing events (120, 200, 765). Two studies suggest that anticholinesterase medications can improve obstructive sleep apnea. Physostigmine reduced the AHI by 20% and improved oxygenation (234a). Donepezil, a second anticholinesterase inhibitor, was recently shown to substantially reduce obstructive apneas and oxyhemoglobin desaturation time in individuals with Alzheimer’s disease (420). The effect was most pronounced in individuals with severe sleep apnea. Whether this cholinergic therapeutic potential is replicated and whether this will be effective in all individuals with obstructive sleep apnea should now be examined.

H) Hypocretin (orexin-A). Under specific circumstances, orexin plays a critical role in motor control, as evidenced by cataplexy and increased sleep paralysis in patients with narcolepsy (642). Orexinergic neurons project to both trigeminal and hypoglossal motoneuronal soma and dendrites (180), and these neurons, like the monoaminergic groups, have reduced c-fos activation in NREM and REM sleep (157); however, like ACh, levels of orexin in motor nuclei are greatest in wake and REM sleep (308). Peever et al. (494) tested the effects of hypocretin-1 and -2 administered into the motor trigeminal nucleus and hypoglossal nucleus on masseter and genioglossus muscle activity, respectively. Both peptides increased masseter electromyographic activity by a similar magnitude, but only hypocretin-1 increased genioglossus activity. The effect of hypocretin-1 was markedly longer lasting. Blocking glutamate transmission attenuates the excitatory ef-
fect of hypocretin in motor nuclei (494). Thus reductions in orexin in upper airway motor nuclei in NREM sleep could contribute to the suppression of upper airway dilator activity in NREM sleep, but this is not expected to contribute to REM sleep atonia in these muscles.

1) LESSER-STUDIED NEUROPEPTIDES. In addition to the above neurotransmitters and neuromodulators, many other neuropeptides modulate brain stem motoneuron activity. Several of these neuromodulators have significant excitatory effects. Vasopressin binds to the V1A receptor on facial and hypoglossal motoneurons and depolarizes the membrane (552). This neuropeptide may play a greater role in newborn and young animals, as the receptor density declines with age (685). Substance P binding to the NK-I receptor is evident in the hypoglossal nucleus, where binding sites decline with intermittent hypoxia (328). Substance P (NK-1) agonist applied to neonatal hypoglossal motoneurons in slice increases the phasic amplitude (747). Gatti et al. (189) have shown that substance P terminals appose the protrusor motoneurons within the hypoglossal nucleus. Oxytocin binding sites are present on hypoglossal motoneurons (373), but the physiological significance of these sites has not been established. Histamine is another excitatory neuromodulator with increased neurotransmission in waking. Thus there remain many targets for potential pharmacotherapeutics for obstructive sleep apnea.

E. Clinical Pharmacotherapeutic Trials for Sleep Apnea

Presently, universally effective drug therapy for sleep apnea deludes us. Overall, drugs with serotonin influences have been emphasized in clinical trials. Both serotonergic and serotonin receptor antagonist drugs have been tested for effectiveness in reducing obstructive sleep-disordered breathing, including L-tryptophan, fluoxetine, paroxetine, trazodone, and mirtazapine. In summary, none of the drugs tested has proven universally effective (52, 222, 313, 573, 590). In general, the sample sizes have been too small to prove or disprove the effects of the drug. In addition, there may be subsets of individuals who will respond to serotonergics, particularly patients with REM sleep predominant events, where a serotonergic antagonist may reduce REM sleep time and phasic activity. In light of hypoglossal excitation with 5-HT1B antagonists, effectiveness of 5-HT1B antagonism should be explored. Three drugs with noradrenergic effects have been tested: protriptyline, amoxetine, and clonidine. The overall findings parallel findings with serotonin drugs: no major effect other than REM sleep suppression (37, 222, 273). Individuals with mild NREM sleep apnea may provide a more promising subset of individuals to study. Muscarinic antagonists have yet to be explored. Similarly, genetic therapies that alter glycine receptor function for pharyngeal motoneuron may be promising. The advent of more accurate in-home polysomnographic devices allowing repeated tests in each subject will advance pharmacotherapeutics for obstructive sleep apnea by allowing adequately powered, multidose, repeated trials. As mentioned previously, because the obstruction is present only in sleep, these apneic events should be readily amenable to drug therapies, provided we identify receptors that will not also have prohibitive side effects.

F. Future Directions for the Neurobiology of Upper Airway Control and for the Development of Pharmacotherapies for Obstructive Sleep Apnea

As stated in the beginning of this section, there are likely orphan G protein-coupled receptors that to date have not been identified for brain stem motoneurons and that may not have the widespread brain activation issues present for established excitatory neurochemicals. Identification of these receptors is a tremendously labor intensive endeavor. A more prudent approach would be to complete the testing of promising serotonergic directions. In particular, adults with mild NREM sleep OSAS may benefit from serotonergic therapies that reduce sleep fragmentation (such as 5-HT2C antagonist therapies), and individuals with REM sleep OSAS and without significant oxyhemoglobin desaturations may benefit from a serotonergic therapy that suppresses REM sleep time. The receptor subtypes that seem promising and worthy of further pursuit include 5-HT1B antagonism and/or 5-HT1A, 5-HT4, or 5-HT7 agonists. The latter subtypes may also act on central respiratory neurons and increase drive to pump muscles. It is possible that combination therapies will be helpful, using drugs that increase serotonergic or other excitatory effects at motoneurons with drugs that suppress the vagal inhibitory effects of ventilation. An animal model of upper airway collapse in sleep is needed to expedite identification of safe and effective serotonergic pharmacotherapeutics. Having dissected the rat hyoid arch and its pharyngeal musculature, we believe that such a model can be developed. Present in the rat, but not human, is a series of connected lateral hyoid arch bones that provide tremendous lateral support for the hypopharynx. We predict that removal of the lateral hyoid bones in obese rats, rabbits, or guinea pigs would increase collapsibility of the upper airway and provide animal models with which to test neuromodulator pathways. With successful therapies substantiated in multiple species, large-scale clinical trials with adequate power and sufficient dose range across well-characterized subsets of individuals with OSAS will be able to determine precisely who may benefit from serotonergic and other pharmacotherapies.
V. CARDIOVASCULAR SEQUELAE OF SLEEP APNEA

A. Introduction

Sleep-disordered breathing (SDB) is recognized as a risk factor for the development of hypertension and other cardiovascular diseases. Because of the associations between SDB and obesity and advancing age, the public health burden of SDB-related cardiovascular disease is expected to rise in the coming years. Fortunately, numerous treatment studies suggest that SDB can be added to the list of "modifiable" risk factors. Although treatment of SDB would seem to be the simplest way to reduce cardiovascular risk, many patients refuse or underutilize the cumbersome state-of-the-art treatments and many remain undiagnosed. In the following section, we discuss current concepts of the mechanisms by which SDB contributes to cardiovascular morbidity and mortality. Progress in this area of investigation will ensure a rational approach to prevention and treatment of the cardiovascular sequelae of SDB.

B. Acute Effects of Sleep Apnea on the Cardiovascular System

1. Central hemodynamic effects

Episodes of OSA produce arterial oxygen desaturation, hypercapnia, intrathoracic pressure oscillations, and in most cases, sleep disruption (Fig. 12). The highly negative intrathoracic pressures generated during obstructed inspiratory efforts produce transient decreases in left ventricular stroke volume (612, 683). Inspiratory strains also produce small, transient reductions in systemic arterial pressure. Cardiac output falls during obstructive apnea, secondary to decreased stroke volume and also to reductions in heart rate, which can be marked in some individuals (96, 187, 592, 635).

Upon resumption of breathing, apnea-induced constraints on stroke volume and heart rate are abruptly removed, allowing release of the augmented cardiac output into a peripheral vascular bed that has been constricted by an increase in sympathetic vasomotor outflow. As a result, the immediate postapnea period is characterized by a marked, transient increase in systemic arterial pressure. This pressor response is caused by sympathetic nervous system activation, because it can be abolished with ganglionic blockade (296, 455, 592). Studies using supplemental oxygen have shown that stimulation of the carotid chemoreceptor by asphyxia (combined hypoxia and hypercapnia) is the most important cause of apnea-induced sympathoexcitation and blood pressure elevation (349, 424). In contrast, the mechanical influence of negative intrathoracic pressure plays little or no role in causing the sympathetically mediated pressor response to apnea (424, 720).

2. Peripheral circulation

Apnea-induced vasoconstriction has been observed in the forearm (13, 264) and the finger (454) of patients with OSA. These findings are surprising because acute
exposure to both hypoxia and hypercapnia causes vasodilation in most vascular beds (1, 540), and they suggest that repeated asphyxic exposures, over time, produce alterations in basic mechanisms of neural and local control of vascular resistance. In support of this notion, time-dependent impairments in hypoxic vasodilation have been observed in healthy humans (192) and rats (390) after exposure to hypoxia.

3. Cerebral circulation

The cerebral circulation is exquisitely sensitive to changes in \( \text{PaO}_2 \) and \( \text{PaCO}_2 \); therefore, episodes of OSA have profound effects on flow in this vascular bed. Cerebral blood flow increases progressively during apneas, followed by abrupt decreases in the postapnea hyperventilation period (31, 217, 309). This oscillatory pattern in cerebral blood flow is determined mainly by fluctuations in \( \text{PaCO}_2 \) with a smaller contribution from apnea-induced increases in arterial pressure (532). These cerebral blood flow oscillations, through their influence on washout of \( \text{CO}_2 \) from central chemoreceptors, may produce breathing instability during sleep (738) (see sect. III).

4. Pulmonary circulation

During episodes of OSA, oscillations in \( \text{PaO}_2 \) produce a cyclical pattern of vasoconstrictions and relaxations in the pulmonary circulation that cause marked fluctuations in pulmonary artery pressure (447, 593). These fluctuations are caused by the local vascular effects of alveolar hypoxia and hypoxemia, because they can be abolished by supplemental oxygen (592).

5. Cardiovascular effects of arousal

Most episodes of OSA are accompanied by arousal from sleep. Sleep disruption per se increases sympathetic nerve activity and blood pressure (422), and arousal appears to augment the pressor effects of asphyxia during SDB (423). In experimental animals, “respiratory” arousals caused by upper airway occlusion produce a pressor response accompanied by vasoconstriction in the hindlimb, whereas “nonrespiratory” arousals caused by acoustic or tactile stimulation elicit a smaller pressor response and hindlimb vasodilation (341). Thus it is likely that alterations in sleep state contribute synergistically to sympathetic vasoconstriction and acute blood pressure elevation caused by apnea. In humans, auditory arousals cause transient perturbations in cerebral blood flow that are dependent on the prearousal sleep state (32).

C. Associations Between Sleep Apnea and Cardiovascular Disease

Case control and epidemiological studies indicate that chronic exposure to OSA plays a pathogenetic role in cardiovascular disease. In the following paragraphs we review the evidence linking OSA and specific disease entities. Putative mechanisms will be discussed in a subsequent section.

1. Hypertension

Case-control (168, 725) and cross-sectional (57, 446) studies reveal an association between OSA and hypertension; however, longitudinal data from the Wisconsin Sleep Cohort provide the most compelling evidence for a causal relationship (502). This study demonstrated a dose-response relationship between OSA severity and incident hypertension, with elevated odds ratios for subjects with AHI of 5–15 and those with AHI >15. In contrast to these findings, a larger prospective study of hypertension incidence did not find a dose-response relationship between AHI and incident hypertension that was independent of obesity (452).

In both population-based studies, the percentage of subjects in the highest AHI category was small (7 and 4%, respectively), and the severity of SDB was not quantified using indexes of nocturnal hypoxemia, which may predict hypertension risk more accurately than AHI does. Obesity is likely to be an important covariate in correlations based on hypoxemia because SDB events of a given duration would cause greater desaturations in overweight versus normal weight individuals secondary to high metabolic rates and low lung volumes (see sect. III).

The initial experimental support for a causal link between OSA and hypertension came from animal models in which exposure to intermittent hypoxia or intermittent airway occlusion equivalent to severe SDB produced blood pressure elevation that was evident, not only during the exposure period, but also when the animals were normoxic and unperturbed (73, 171). This rise in blood pressure, which was dependent on activation of the sympathetic nervous system by carotid chemoreceptors, was caused mainly by intermittent hypoxia. The addition of intermittent hypercapnia did not augment the hypertensive effect (348), and experimental paradigms that employed repetitive arousals from sleep without hypoxia did not result in blood pressure elevation (33, 73).

More recently, investigators have used chronic intermittent hypoxia (CIH) paradigms that model “mild” (244) and “moderate” (9, 292, 315, 390, 686) levels of SDB. Elevations in blood pressure were observed following most (9, 292, 390, 686) but not all (315) of the studies that employed moderate-level exposures (15–20 events/h). “Mild” CIH (10 events/h) caused a more modest increase in blood pressure in male rats and ovariectomized female rats and only a slight increase in intact females (244).

Reductions in daytime blood pressure following treatment of OSA with nasal CPAP provide further strong evidence for a causal relationship. Some investigators have observed statistically significant reductions in 24-h...
mean blood pressures following CPAP (Fig. 13) (41, 158, 237, 393, 451, 503), whereas others have not (81, 560). A potential reason for this inconsistency and for the modest blood pressure-lowering effects of CPAP in some of these papers is that many of the subjects were normotensive; CPAP would be expected to have minimal impact in such individuals. In contrast, relatively large CPAP effects on 24-h blood pressure have been demonstrated in hypertensive patients (237), particularly those with resistant hypertension (393).

The fact that 24-h mean blood pressure does not decrease with CPAP in all patients may be due to inconsistent compliance with treatment, differences in treatment duration or length of exposure to OSA, and the possibility that OSA-related vascular remodeling is irreversible, or it may reflect the multifactorial nature of hypertension. It has recently been suggested that CPAP treatment reduces 24-h mean blood pressures primarily in patients with excessive daytime sleepiness (35, 311, 560). In a multiple regression analysis, baseline blood pressure, improvement in Epworth sleepiness score, BMI, decrease in heart rate, and decrease in time spent at <90% saturation (but not AHI) emerged as important predictors of CPAP-induced decrease in 24-h blood pressure (558).

The effects of CPAP treatment on nighttime blood pressure are unequivocal. Episodes of SDB, even mild apneas and hypopneas, result in substantial acute blood pressure elevations (423); therefore, it is not surprising that elimination of these events lowers average nocturnal values. Numerous investigators have reported CPAP-induced decreases in both systolic and diastolic pressure during sleep and restoration of the nocturnal “dipping” pattern (393). Restoration of this normal circadian blood pressure pattern is an important benefit of CPAP because daytime cardiovascular events and complications are more prevalent in individuals who lack sleep-related falls in blood pressure (134, 697). Moreover, elimination of OSA-related surges in left ventricular afterload may improve cardiac remodeling (140). Thus the benefits of CPAP therapy on cardiovascular risk may extend well beyond demonstrable effects on 24-h mean blood pressure.

The blood pressure-lowering effects of alternative therapies for OSA have recently begun to be tested. In one study, CPAP significantly lowered 24-h blood pressure, whereas supplemental oxygen did not (451). Treatment with dental appliances caused modest yet statistically significant reductions in 24-h mean blood pressure (201), whereas in another study there was no change in blood pressure with any treatment (CPAP, oral device, or placebo pill) (36). It is important to note that in both studies, most of the patients were normotensive or were receiving antihypertensive medications.

The effect of OSA on blood pressure in children and adolescents has received relatively little study. In a clinic-based study, Amin et al. (12) found that 24-h mean blood pressures did not differ according to OSA status; however, children with OSA had smaller nocturnal declines and greater blood pressure variability during sleep and wakefulness. Leung et al. (350) observed that OSA and hypertension were correlated only in obese children. Several recent population-based studies demonstrated that children with SDB (AHI >5) have higher blood pressures versus control subjects, even after adjustment for obesity (56, 155, 354). Alleviation of OSA in children may prevent hypertension; however, at this time no treatment data are available.

In summary, a causal link between severe OSA and hypertension has been firmly established via animal models (73, 171, 390) and CPAP treatment studies that have shown, in hypertensive patients with severe OSA, that blood pressure decreases when SDB is eliminated (41, 503). Whether mild SDB also raises blood pressure is less clear. Multiple epidemiological studies have reported a dose-response relationship between OSA severity, as indicated by AHI, and presence of hypertension (57, 141, 446, 502). In contrast, a larger, more recent prospective study failed to find a statistically significant dose-response relationship that was independent of obesity.

![Fig. 13. Mean arterial pressures in patients with OSA before and after effective CPAP (A) and subtherapeutic CPAP (B). [From Becker et al. (41).]](http://physrev.physiology.org/doi/10.220.32.247)
This discrepancy may be due, at least in part, to demographic differences among these cohorts (e.g., older subjects in the recent study; Ref. 452). A more important reason for inconsistency in these findings regarding the dose-response relationship between SDB and hypertension may be that all but one (446) of these previous studies used AHI as a measure of SDB severity. AHI and other measures of event frequency are inadequate indicators of the amount of intermittent hypoxia, which is likely to be the major cause of cardiovascular dysfunction and disease in the setting of SDB. Furthermore, it may not be possible to firmly establish a dose-response relationship between OSA and hypertension via population-based studies because of the measurement error inherent in such studies (e.g., low reproducibility of single overnight sleep studies and/or measurements of blood pressure, indirect measures of breathing). Confirmation of the dose-response relationship will require prospective studies of the hypertensive effects of mild, moderate, and severe SDB in experimental models and clinical intervention studies in patients across all levels of AHI.

It is commonly believed that more than half of all patients with severe OSA are hypertensive (168, 508, 616, 725). Why do some individuals with OSA develop hypertension while others do not? We believe that the answer lies in the multifactorial nature of this complex disease; OSA is but one of several known risk factors for hypertension. Some individuals may be at increased risk for OSA-related hypertension on the basis of their genetic make-up and/or presence of other prohypertensive characteristics, whereas others may have relative protection from the adverse effects of OSA. An important challenge for future researchers is to identify the patients most at risk for hypertension and other cardiovascular sequelae.

1. **Left ventricular dysfunction**

Individuals with SDB are more likely to have CHF than those without SDB (608). Conversely, the presence of SDB in patients with CHF is much higher than in the population at large (282). Although central sleep apnea is common in CHF, a significant fraction of patients also experiences obstructive apneas and hypopneas (282, 617) (see sect. III).

Independent associations between OSA and more subtle impairments in ventricular function have been difficult to document. In patients with severe OSA without coexisting heart disease, left ventricular ejection fraction was only modestly reduced relative to control subjects (53 vs. 61%) (7). Ejection fractions below 50% were noted in only 8% of patients with moderate to severe OSA (mean AHI, 47 events/h) (326). Diastolic dysfunction was observed in approximately one-third of patients with severe OSA; however, no comparison group was studied (179). A recent study of obese patients with severe OSA and BMI-matched control subjects revealed depressed diastolic function and increased left atrial volume in OSA patients, whereas left ventricular mass was comparable in the two groups (475). In contrast, several other studies have shown that OSA is associated with left ventricular hypertrophy, even in the absence of systemic hypertension (140, 232, 450). In a population-based study, an association was observed between SDB (AHI >15) and left ventricular hypertrophy using Cornell voltage criteria (87), but not classic voltage criteria (245).

Interestingly, other investigators have not observed decreases in left ventricular systolic function, diastolic function, or increased mass in OSA patients (219, 448). The reasons for this discrepancy are not obvious; however, in one study, SDB was not completely absent in the control subjects (they were nonapneic snorers) (219). In the other study, the expected associations between BMI and hypertension and left ventricular mass were seen, but no independent effect of OSA could be discerned (448).

Nevertheless, studies in experimental animals support the notion that OSA can negatively affect ventricular function. In dogs, several weeks’ exposure to severe, repetitive airway obstructions during sleep increased blood pressure, caused left ventricular hypertrophy, and reduced ejection fraction (485). Increased left ventricular mass has been observed in rats exposed to CIH (169).

Limited additional evidence for a causal relationship between OSA and ventricular dysfunction comes from observations of improved heart function following CPAP. In patients with systolic dysfunction and moderate to severe OSA, CPAP treatment improved ejection fraction, functional status, and quality of life (293, 387). However, because neither trial included a placebo treatment arm, these results must be interpreted with caution (478).

Strong, consistent evidence for OSA as a cause of ventricular dysfunction is not available; however, the large increases in ventricular afterload produced by acute episodes of OSA are likely to negatively impact individuals with existing ventricular dysfunction. OSA is often present in patients with severe ventricular dysfunction; however, in many of these individuals CHF is the cause, not the consequence, of OSA.

2. **Stroke**

An association between OSA and stroke has been observed in numerous cross-sectional studies (17, 39, 143, 198, 413, 487, 608, 716); however, in most cases it was not possible to determine whether OSA preceded the onset of stroke and thus could be involved in pathogenesis. Arzt et al. (17) recently performed a longitudinal analysis of the association between SDB and stroke risk. They found that moderate to severe SDB (AHI ≥20) was associated with increased risk of incident stroke, whereas no increased risk was observed in subjects with mild SDB. The
increased risk of stroke associated with SDB appeared to be partially independent of hypertension and confounded by obesity.

Yaggi et al. (744) reported the incidence of stroke (including transient ischemic attack) or death from any cause in individuals with previously diagnosed OSA. They found a significant association between SDB, defined as AHI ≥5, and stroke or death from any cause that persisted after statistical adjustment for confounding factors. The authors reported a dose-response relationship between SDB severity and risk; however, because the combined outcome measure of stroke or death from any cause was used, it is not possible to ascertain the specific effects of OSA on risk of stroke. Munoz et al. (433) conducted a population-based study of OSA and stroke risk in the elderly. They found that incident stroke was nearly three times as likely in subjects with AHI ≥30 versus those with AHI <30. The risk of stroke and transient ischemic attack was assessed prospectively in patients with verified coronary artery disease (417). In this group, OSA (AHI >10) was independently associated with incidence of these cerebrovascular end points. A large clinic-based study assessed the effect of OSA on risk of all-cause mortality in patients with preexisting stroke (568). The odds ratio for death from any cause was higher in patients with OSA (mean AHI ~35) than in those without OSA (AHI <15). Interestingly, central sleep apnea was not associated with increased risk.

Viewed collectively, these studies demonstrate an independent association between moderate to severe (AHI >30 events/h), but not mild, OSA and risk of stroke. Treatment of OSA appears to decrease the risk of stroke, because the incidence of fatal and nonfatal cardiovascular events is similar in CPAP-treated individuals and control subjects (391). Interestingly, the association between OSA and stroke risk is partially independent of hypertension (17, 744), an established major risk factor for stroke, which suggests that additional pathogenetic mechanisms are operant (see below).

4. Coronary artery disease

Case-control studies have reported a high prevalence of SDB in individuals with clinically verified coronary artery disease (417–419, 495). In these studies, the odds ratios for OSA as an independent predictor of coronary disease approximated those of known risk factors such as diabetes mellitus, hypertension, and obesity. In contrast, large population-based studies have not detected a strong independent association between SDB and coronary disease (245, 608). One possible explanation for divergent findings in clinic- versus population-based studies is differences in the severity of coronary artery disease. All subjects in the former studies had clinically manifest coronary artery disease, whereas in the population stud-

ies, subclinical coronary disease may not have been recognized.

Coronary artery calcification, an indicator of subclinical atherosclerosis, was quantified by electron-beam computed tomography in patients referred to a sleep clinic, none of whom had signs or symptoms of coronary artery disease (638). The authors reported a dose-response relationship between OSA severity and coronary artery calcification; however, the odds ratio for calcification was statistically distinct from the reference group only in individuals with severe OSA.

Regardless of whether OSA plays a causal role in the development of coronary atherosclerosis, acute episodes of OSA may influence morbidity and mortality in patients with coexisting coronary disease. In many such individuals, apneas are accompanied by nocturnal angina pectoris and electrocardiographic evidence of myocardial ischemia (2, 176, 220), which can be ameliorated with nasal CPAP (176, 220, 496). Apnea-induced hemodynamic perturbations may have deleterious effects on unstable coronary lesions, leading to plaque rupture and thrombogenesis.

5. Cardiac arrhythmias

Apneas produce multiple arrhythmogenic factors: alterations in cardiac sympathetic and parasympathetic activities, myocardial hypoxemia (585), and intrathoracic pressure fluctuations that deform and alter the size of the cardiac chambers (108). In the initial report on this topic, arrhythmias (most commonly bradyarrhythmias, premature ventricular contractions, and atrial fibrillation/flutter) were detected by 24-h ambulatory monitoring in nearly 50% of patients with severe OSA (212). Interestingly, premature ventricular contractions were more common during wakefulness than sleep. Sleep-related bradyarrhythmias and atrial fibrillation/flutter were abolished after treatment, whereas the premature ventricular contractions persisted. In a large, population-based study, the prevalence of complex ventricular ectopy and atrial fibrillation was greater in subjects with severe SDB (≥30 events/h) versus those without SDB (402). In another recent report, patients referred to a cardiology clinic for atrial fibrillation were twice as likely to have OSA, as assessed by the Berlin questionnaire, than patients referred for management of other cardiovascular disease (184). An increased rate of recurrence of atrial fibrillation after cardioversion has been observed in patients with untreated versus treated OSA (291).

Significant associations were observed between presence of bradyarrhythmias and severity of SDB (462). All observed nocturnal bradyarrhythmias (e.g., sinus pause, AV block) occurred during apneas, whereas only about one-third of tachyarrhythmias were temporally related to SDB events (462). In several case series, treatment of OSA

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reduced the occurrence of nocturnal cardiac rhythm disturbances (40, 207, 223, 310, 680).

Taken together, these findings indicate that patients with moderate to severe OSA are at increased risk for arrhythmias, especially bradycardias, during sleep. While the clinical significance of OSA-related arrhythmias is not clear, an obvious concern is their potential contribution to sudden cardiac death. Gami et al. (183) reported that OSA causes a shift in the diurnal pattern of sudden cardiac death occurrence from the early morning waking hours (6:00 a.m. to noon) to the sleeping hours (midnight to 6:00 a.m.) (183).

6. Pulmonary hypertension

Episodes of OSA cause marked, acute elevations in pulmonary artery pressure (see above). Over time, structure and function of the pulmonary circulation are altered, resulting in fixed elevations in pressure. CIH causes sustained increases in pulmonary artery pressure in mice (159). Increased prevalence of daytime pulmonary hypertension has been observed in patients with OSA, with estimates ranging from <20 to >70% (26, 90, 174, 318, 325, 330, 570, 572, 577, 713). This wide range of estimates is due in part to methodological differences (i.e., right heart catheterization versus Doppler echocardiography). The independent contribution of OSA to diurnal pulmonary hypertension is impossible to judge from some of these studies, because subjects with coexisting lung and heart disease were not excluded. In studies where such subjects were excluded, the prevalence of OSA-related pulmonary hypertension has been estimated at ~20% (6, 26, 90, 577). The incidence of pulmonary hypertension in patients with OSA is unknown, because population-based longitudinal data have not been reported.

If every apnea of sufficient duration causes pulmonary vasoconstriction, why isn’t fixed daytime pulmonary hypertension a universal finding in patients with OSA? Interindividual differences in hypoxic sensitivity of the pulmonary vasculature (435) probably play a role, as do the frequency and severity of nocturnal desaturations and presence of comorbid conditions. In OSA patients without clinically significant heart and lung disease, pulmonary artery pressure was positively correlated with age, BMI, percentage of total sleep time with \( S_{\alpha O_2} \leq 90\% \) (but not AHI), and daytime \( P_{aCO_2} \) (6, 26), and negatively correlated with spirometric measures of pulmonary function (26) and daytime \( P_{aO_2} \) (6, 26). These findings suggest that even subtle changes in pulmonary function, in the absence of frank lung disease, can contribute to the development of pulmonary hypertension in patients with OSA. It is important to note, however, that pulmonary hypertension could also be a cause of abnormal arterial blood gases during wakefulness because of its adverse effect on ventilation-perfusion distribution. In addition, left ventricular hypertrophy (232) and diastolic dysfunction (179), which are prevalent in patients with OSA, could cause postcapillary pulmonary arterial hypertension in susceptible individuals.

Even though daytime pulmonary artery pressures may not reach the hypertensive range in all patients with OSA, they are higher than in matched control subjects (6, 16). These elevations are mild relative to those observed in primary pulmonary hypertension, and their clinical significance is unknown. In some patients with OSA, pulmonary artery pressures are within normal limits at rest but increase to hypertensive levels during exercise (242, 521, 679), suggesting that capillary recruitment during exercise, a major mechanism for keeping pulmonary artery pressure low in the face of increasing cardiac output, is limited. A possible explanation for this limitation is a reduction in the total number of vessels in the pulmonary vascular bed (rarefaction), similar to that responsible for capillary pulmonary hypertension in chronic obstructive pulmonary disease (247, 566). OSA-related pulmonary hypertension may be responsible, at least in part, for exertional dyspnea and exercise intolerance in affected individuals (363).

Fixed increases in pulmonary artery pressure caused by OSA may have clinical significance. Increased right ventricular mass and decreased ejection fraction have been observed in OSA patients, particularly those with diurnal arterial blood gas abnormalities (67, 174, 577). The long-term consequences of OSA-related pulmonary hypertension are not known; however, in patients with chronic pulmonary disease, pulmonary hypertension is associated with increased morbidity and mortality (381, 382, 566, 604). Following CPAP treatment, decreases in daytime pulmonary artery pressures have been observed, regardless of whether the diagnosis of pulmonary hypertension was present (6, 16, 571).

7. Summary

Numerous epidemiological and case-control studies have demonstrated statistically significant associations between SDB and cardiovascular disease. In these studies the odds ratios were highest for individuals with moderate to severe SDB; however, dose-response relationships were rarely apparent. Hypopneas that produced >4% desaturation were independently associated with coronary artery disease, whereas those with <4% desaturation were not (533). Viewed collectively, these studies suggest that a threshold level of SDB (~25–30 events/h of sleep) and significant oxygen desaturation are required to produce cardiovascular disease.

Is OSA-related cardiovascular risk modifiable? Recent nonrandomized long-term treatment studies suggest that it is. Fatal and nonfatal cardiovascular events were less frequent in treated versus untreated patients with severe OSA (391). Other investigators reported a treat-
ment-related enhancement of event-free survival, even in people with mild to moderate OSA; however, in this study, follow-up durations were different in treated and untreated patients, compliance with other therapies (i.e., medications) was not documented, and a substantial fraction of subjects was lost to follow-up (77). Therefore, further study is required before definite conclusions can be reached regarding the effects of treatment on cardiovascular risk in mild to moderate OSA.

D. Pathophysiological Links Between Sleep-Disordered Breathing and Cardiovascular Disease

OSA alters many aspects of cardiovascular structure and function via multiple pathways; however, when the existing evidence is viewed collectively, several common themes emerge. In subsequent paragraphs we highlight the roles played by neurohumoral activation, oxidative stress, and inflammation in OSA-induced cardiovascular morbidity and mortality.

1. Alterations in neurohumoral control of the circulation

Sympathetic nervous system activity is heightened in OSA patients during sleep and during wakefulness (83, 233, 439). Reductions in muscle sympathetic nerve activity (MSNA) have been documented following CPAP treatment (231, 265, 438, 707), indicating a causal relationship between OSA and sympathetic activation.

In healthy humans, brief exposures to continuous or intermittent asphyxia during wakefulness cause increases in MSNA that persist after reestablishment of normoxic, normocapnic conditions (Fig. 14) (113, 421, 736). Long-lasting sympathoexcitation caused by asphyxia is critically dependent on hypoxia, rather than generic chemoreflex stimulation, because it does not occur following hypocapnia alone (741). The mechanisms that maintain high levels of MSNA after withdrawal of chemical stimuli are not known; however, available evidence suggests that this long-lasting sympathoexcitation has both reflex and central nervous system origins.

In patients with OSA, plasma levels of angiotensin II (ANG II) and aldosterone are elevated, and CPAP treatment causes decreases in plasma renin and ANG II that correlate with reductions in blood pressure (414). The blood pressure elevation produced by CIH in rats can be prevented by blockade of ANG II type I receptors (AT_1R), by suppression of the renin-angiotensin-aldosterone system with high-salt diet, and by renal nerve denervation (166, 173). The latter finding suggests an important role for circulating ANG II; however, production of ANG II in

![Fig. 14](http://physrev.physiology.org/)

**Fig. 14.** In healthy humans, brief (20-min) exposure to intermittent asphyxia causes sympathetic activation that persists after normalization of blood gases, not only during the interasphyxia phases, but also in the room air recovery period. [From Xie et al. (736).]
the vascular wall and in other tissues may also contribute importantly to the blood pressure raising effect of CIH.

2. Augmentation of carotid chemoreflex function

In rats, CIH increases diurnal blood pressure, but only when the sympathetic nervous system and the carotid chemoreceptors are intact (170, 171). CIH exposure also increases basal sympathetic outflow and results in enhanced sympathetic activation during subsequent acute hypoxic exposures (131, 206, 389, 529, 615). Moreover, CIH enhances carotid body sensory activity, as evidenced by increased rates of carotid sinus nerve discharge during normoxia and upon reexposure to hypoxia (498, 551, 615). These findings, which indicate sensitization of the carotid chemoreceptor, are consistent with human data showing that 1) supplemental oxygen reduces sympathetic outflow in OSA patients but not control subjects (439), and 2) hypoxia-induced sympathetic activation is enhanced in OSA patients versus controls (440).

Reactive oxygen species (ROS) generated during hypoxia-reoxygenation cycles play a role in the carotid chemoreflex sensitization caused by CIH. Pretreatment with a superoxide anion scavenger prevented CIH-induced sensory long-term facilitation of the carotid body (498) and long-term facilitation of respiratory motor output (499). The mitochondrion appears to be a source of ROS in this model (322, 528); however, oxidases in the plasma membrane and cytoplasm (e.g., NADPH oxidase and xanthine oxidase) may also contribute importantly. Several lines of evidence suggest that ANG II-induced activation of NADPH oxidase and subsequent ROS production may contribute importantly to the chemoreflex sensitization produced by CIH. Infusion of ANG II into the isolated carotid body increases carotid sinus nerve activity (336), and exposure to hypoxia upregulates AT1R in the carotid body (335). Systemic infusion of ANG II increases carotid sinus nerve discharge and augments the amount of sympathoexcitation elicited by acute exposure to hypoxia (360). Both of these effects can be abolished by concomitant infusion of an AT1R antagonist, a superoxide dismutase mimetic, or an inhibitor of NADPH oxidase. Taken together, these findings indicate that ANG II sensitizes the carotid chemoreflex via AT1R activation and NADPH oxidase-derived oxidative stress (359, 360). Exposure of rats to CIH augments chemoreflex control of sympathetic outflow and results in increased AT1R expression and increased superoxide production in carotid body, all of which can be prevented by in vivo treatment with losartan, an AT1R antagonist (389). A potential stimulus for upregulation of carotid body ANG II/AT1R by CIH is an altered expression of oxygen-sensitive potassium channels (288).

Endothelin-1 (ET-1), which is present in glomus cells of the carotid body, produces chemoxecitation by binding to ET-1A receptors (94, 527). In cats, 4 days of CIH produced a 10-fold increase in the expression of ET-1 receptors in the carotid body, increased basal carotid sinus nerve discharge, and an augmentation in the response to acute hypoxia (551). This potentiation, which could be abolished by the endothelin receptor antagonist bosentan, was larger in perfused versus superfused isolated carotid bodies, which suggests that the excitatory effect of ET-1 is mediated, at least in part, by its vasoconstrictor effect.

Intermittent (but not continuous) application of 5-HT elicits sensory long-term facilitation of the isolated carotid body of rats and mice (500). Although 5-HT is a potent vasoconstrictor, this adaptation was not dependent on vascular responses because it occurred in the superfused carotid body. This sensory long-term facilitation could be prevented by concomitant application of ketanserin (5-HT1 receptor antagonist), apocynin (an inhibitor of NADPH oxidase), and N-acetylcysteine (an antioxidant).

In contrast to ANG II, ET-1, and 5-HT, nitric oxide (NO) is an inhibitor of carotid body chemosensitivity (656). In rabbits, intravenous administration of Nω-nitro-L-arginine (L-NNA), a NO synthase inhibitor, increased the basal rate of carotid sinus nerve discharge and enhanced the response to hypoxia (656). Conversely, administration of L-arginine, the substrate for NO synthase, decreased baseline discharge and attenuated hypoxic chemosensitivity (656). The carotid body contains two isoforms of NO synthase: eNOS expressed in blood vessels and nNOS expressed in ganglion cells (526). In rats, exposure to CIH causes downregulation of nNOS mRNA (712) and protein (389) in carotid body; thus removal of the inhibitory influence of NO may be responsible, at least in part, for CIH-induced increases in carotid chemoreflex sensitivity.

3. Decrements in baroreflex function

Depressed baroreflex control of heart rate has been documented in patients with OSA (28, 84, 449), and in one study, impaired baroreflex control of MSNA was also observed (84). However, patients and control subjects were not always matched on the basis of hypertension, which can per se alter the set-point and sensitivity of sinoaortic baroreceptors (91). In matched OSA patients and control subjects, phenylephrine-induced increases in blood pressure elicited comparable heart rate and MSNA responses (439). In the same study, nitroprusside-induced decreases in blood pressure revealed OSA-related impairment of the sympathetic, but not the heart rate, component of the baroreflex response (439). When hypertensive and normotensive OSA patients were compared, decreased baroreflex control of heart rate was observed in hypertensive but not normotensive patients (28, 769). Taken together, these findings suggest that depressed
Baroreflex function in OSA is attributable, at least in part, to coexisting hypertension. Baroreflex sensitivity has been studied in experimental models of OSA. In healthy humans, 30-min exposure to voluntary, repetitive, end-expiratory apneas failed to alter the gain of baroreflex control of heart rate and MSNA, even though both operating points were shifted toward higher heart rates and higher levels of sympathetic activity (415). A dog model of intermittent airway obstructions during sleep also caused baroreflex resetting, but failed to alter baroreflex sensitivity (72). In rats exposed to CIH, a decrease in baroreflex sensitivity was observed after 2 wk; however, arterial pressure became elevated after only 5 days, which suggests that decreased baroreflex sensitivity was a consequence, not the cause, of the blood pressure elevation (329).

It is not clear whether decreased baroreflex control of heart rate in OSA patients represents a neural adaptation or whether it is secondary to decreased arterial compliance in the carotid sinuses and aortic arch. Several investigators have observed OSA-related increases in arterial stiffness (28, 140, 311, 509, 669, 688). In OSA patients who were treated with CPAP, improvements in baroreflex sensitivity occurred in parallel with decrements in arterial stiffness (449).

It is unlikely that alterations in baroreflex function, per se, are responsible for elevations in diurnal blood pressure in OSA; however, they may result in impaired ability to buffer the pressor responses generated by episodes of apnea. This notion is supported by the observation that the surges in MSNA caused by obstructive apneas are, at least in part, resistant to baroreceptor inhibition (378).

4. Increases in central sympathetic outflow

In addition to effects on chemoreflex function, chronic exposure to intermittent hypoxia may augment central sympathetic outflow. Again, ANG II is a likely contributor to this process. Sympathetic premotor neurons in the brain stem, which are important modulators of postganglionic sympathetic discharge, receive excitatory inputs from higher centers such as the paraventricular nucleus (PVN) of the hypothalamus and the circumventricular organs (115, 185, 541).

In its role as a regulator of central sympathetic outflow, ANG II has important inhibitory interactions with NO (366, 770). CIH exposure decreases nNOS expression in the PVN and increases AT₃R expression in the circumventricular organs (712). Cortical regions that modulate central sympathetic outflow may also be involved in CIH-induced sympathoexcitation. Increased Fos-like immunoreactivity was observed in medial prefrontal and insular cortex following 30 days of CIH (615).

5. Alterations in local vascular regulation

Several lines of evidence indicate that endothelial function is impaired in patients with OSA. Reductions in endothelium-dependent vasodilation in the forearm have been demonstrated by invasive and noninvasive means (110, 295, 312, 445). In both cases, indicators of nocturnal hypoxemia (e.g., minimum arterial oxygen saturation, amount of time with saturation <90%) better predicted the degree of endothelial dysfunction than did the frequency of apneas and hypopneas. A causal relationship between OSA and endothelial dysfunction was demonstrated by a study in which flow-mediated dilation in the forearm was improved by CPAP treatment (270). This beneficial effect was lost when CPAP was temporarily withheld. L-NMMA, a NO synthase inhibitor, caused greater reduction in forearm blood flow after versus before CPAP treatment, which suggests that elimination of OSA augmented resting NO availability (340).

Pulmonary artery responses to ACh, sodium nitroprusside (SNP), and L-NMMA were assessed in OSA patients before and after CPAP treatment (339). The decrease in pulmonary artery blood flow produced by L-NMMA was more pronounced after treatment, suggesting that NO levels in the pulmonary circulation were depressed in the untreated condition. In addition, small increases in ACh-induced vasodilation were observed after treatment (339).

Several other observations suggest that OSA reduces the bioavailability of NO. Decreased plasma levels of NO derivatives and normalization of these levels following CPAP treatment have been observed in patients with OSA (11, 267). Scavenging of NO by ROS is a potential explanation for the decrease in its bioavailability. In patients with OSA, increased production of superoxide by neutrophils (594), increased biomarkers of lipid peroxidation (345), and increased levels of 8-isoprostanes (11, 86) have been observed.

Data from animal models are consistent with these findings of decreased NO bioavailability. In rats exposed to CIH, acetylcholine-induced dilation in cremaster arteries is diminished and the constrictor response to acute NO synthase inhibition is smaller than in control rats (664). CIH also attenuates ACh-induced vasodilation in the cerebral and skeletal muscle circulations, whereas it has no effect on nitroprusside-induced vasodilation (512).

Interestingly, CIH in rats also attenuates norepinephrine-induced vasoconstriction (511). In vivo treatment with a superoxide dismutase mimetic prevented this attenuation, which suggests that the impairment was caused by excess superoxide ion and oxidative/nitrosative stress (511). Other investigators have observed that exposure to intermittent asphyxia enhances ET-1-induced vasoconstriction in the mesenteric circulation (10). In lungs isolated from rats exposed to CIH, vasoconstriction

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produced by a thromboxane mimetic that was augmented (630). In the forearms of patients with OSA, Hedner and colleagues observed impaired norepinephrine-induced vasoconstriction (210) and enhanced ANG II-induced vasoconstriction (314). Thus it appears that the effects of CIH exposure on vasocostructor responsiveness are specific to the vascular bed and vasoactive substance under study.

There is some evidence that ET-1 contributes to CIH-induced vascular dysfunction. ET-1 receptor blockade lowers blood pressure in rats previously exposed to CIH (292), and CIH increases plasma levels of ET-1 (686). Elevated plasma levels of ET-1 are not consistently seen in OSA patients (194, 208, 286, 507, 567, 764); nevertheless, high local concentrations of ET-1 could adversely affect vascular function, as has been shown in the carotid body (551), in the absence of elevated plasma levels.

How does CIH exposure cause oxidative/nitrosative stress in the vascular wall? Recent evidence points to two enzymes, NADPH oxidase and xanthine oxidase, as potential sources of ROS that contribute to CIH-induced vascular dysfunction. Although the mechanism is unclear, exposure to CIH has been shown to enhance NADPH-stimulated superoxide production in rat mesenteric arteries (686). It is possible that CIH-induced activation of the renin-angiotensin system contributes to vascular oxidative stress via the known stimulatory effects of ANG II on NADPH oxidase- (342) and xanthine oxidase-dependent superoxide production (337). Superoxide generated in this manner would be expected to react with NO to form peroxynitrite, thereby reducing NO availability. Peroxynitrite, in turn, can oxidize tetrahydrobiopterin (BH₄), a critical cofactor for NO synthase, causing the enzyme to produce superoxide instead of NO (so-called “uncoupling”) (297). In patients with OSA, flow-mediated dilation in the forearm was enhanced by allopurinol treatment (151), which suggests that xanthine oxidase-derived superoxide plays an important role in OSA-induced endothelial dysfunction.

Reactive oxygen species may also contribute to OSA-induced cardiovascular morbidity via their role as initiators of the inflammatory response (343). Increased levels of C-reactive protein and various proinflammatory cytokines have been reported in OSA patients versus control subjects (564, 609, 699, 749). In a rat model, exposure to intermittent airway obstruction elicited a systemic inflammatory response (434).

The mechanisms by which inflammation contributes to OSA-induced vascular dysfunction are not known; however, recent evidence points to the involvement of the T-lymphocyte. This cell is known to play an important role in ANG II-induced hypertension and endothelial dysfunction via NADPH oxidase-induced superoxide production (216). Furthermore, ET-1 is a potent activator of T-cells (122).

Inflammation may be an important link between increased sympathetic nervous system activity and vascular dysfunction in OSA. Chronically elevated sympathetic activity evokes an inflammatory cascade in several organs and vascular beds (761), and T-cell-rich tissues like the liver and spleen are densely innervated with sympathetic nerves. Moreover, hypoxia-induced increases in renal sympathetic nerve activity activate the renin-angiotensin-aldosterone system. Mineralocorticoid receptor stimulation induces not only inflammation, but also oxidative stress, leading to endothelial dysfunction and vascular remodeling (75). It is therefore tempting to speculate that inflammation is an important link between the neurohumoral and vascular manifestations of OSA in the pathogenesis of hypertension and atherosclerosis.

What is the impact of OSA-induced endothelial dysfunction on vascular regulation? In patients with OSA, attenuated hypoxic vasodilation in the forearm (546a, 550) and the cerebral circulation (175, 546a) have been observed, as have attenuated cerebrovascular responses to hypercapnia (135, 546a). In a large population-based study, decrements in cerebrovascular CO₂ reactivity were associated with measures of nocturnal oxygen saturation, but not AHI, suggesting that the degree of hypoxemia is a more important contributor to these functional impairments than frequency of events (545). These alterations in vascular regulation may negatively impact tissue perfusion during acute episodes of apnea. Moreover, the resulting enhanced pressor responses to apneas may contribute to loss of the normal sleep-related decline in blood pressure (i.e., “nondipping”). The functional consequences of OSA-induced endothelial dysfunction are not well understood; however, OSA-induced impairments in vascular function may compromise blood flow regulation during other stressors, such as exercise.

6. Alterations in arterial wall structure and biomechanics

Increased carotid intima-media thickness (412, 669) and increased arterial stiffness (28, 140, 311, 509, 669, 688) have been observed in individuals with OSA. Blood levels of NO and ET-1, endothelium-derived regulators of vascular stiffness with opposing actions, are decreased and increased, respectively (267, 507). In rats, 14-day exposure to CIH increases vascular wall stiffness in skeletal muscle resistance arteries (511).

Mitogenic factors known to participate in remodeling (e.g., vascular endothelial growth factor, basic fibroblast growth factor, platelet-derived growth factor) are upregulated during hypoxia (85, 142) and during inflammation (75). Inflammation triggers secretion of enzymes that disrupt the balance between matrix metalloproteinases and their inhibitors (274). ANG II (197, 767) and aldosterone (75) are well-established promoters of vascular inflammation.
tion and remodeling; therefore, CIH-induced activation of the renin-angiotensin-aldosterone system may contribute importantly to CIH-induced alterations in vascular structure and biomechanics.

Chronic sympathetic activation may contribute to vascular remodeling via inflammation (761) and/or release of catecholamines that induce vascular wall growth (58). It has recently become evident that adventitial fibroblasts contribute to hypoxia-induced vascular remodeling (648). The release of ATP from adrenergic nerve terminals during hypoxia-induced sympathetic stimulation causes proliferation and migration of adventitial fibroblasts into the intima and media of pulmonary arteries (190) and may also be a stimulus for remodeling in the systemic circulation. In addition, surges in sympathetic outflow during episodes of apnea produce cyclic increases in arterial pressure and blood flow. The cyclical stretch caused by these surges may trigger adaptations in endothelial cells, vascular smooth muscle, and extracellular matrix aimed at normalizing wall stress (689, 727).

Remodeling of the pulmonary circulation has been investigated in animal models of OSA. In mice, CIH raises pulmonary artery pressure (80, 159), produces right ventricular hypertrophy (80, 159), and causes muscularization of pulmonary arteries similar to that caused by continuous hypoxia (159). In rats, CIH-induced right ventricular hypertrophy has been documented in many previous investigations (167, 169, 171, 316, 400, 630). In addition to its effect on NO bioavailability in pulmonary resistance vessels (see above), there is preliminary evidence to suggest that CIH causes pulmonary hypertension via inflammatory pathways (63, 196, 647).

7. Development of atherosclerotic lesions

Independent associations exist between OSA and major risk factors for atherosclerosis, including hypertension (502) as well as insulin resistance (268) and hypercholesterolemia (104) (see sect. vi). Moreover, early signs of atherosclerosis are evident in OSA patients, even in the absence of traditional risk factors, and CPAP treatment results in reversal of these changes (139). In an animal model, CIH has been shown to cause atherosclerotic lesions in mice fed a high-cholesterol diet (583). In this study, marked progression of dyslipidemia and increased serum markers of lipid peroxidation were also observed.

OSA-induced oxidative stress and inflammation, which were discussed in relation to impaired vascular function and structure (above), also are likely contributors to the development of atherosclerotic lesions. High levels of ROS in endothelial, vascular smooth muscle, and adventitial cells are known to initiate atherogenic processes (225). ROS-activated proinflammatory transcription factors, such as activator protein-1 and nuclear factor-κB, stimulate the production of inflammatory cyto-
kines that cause proliferation of vascular smooth muscle cells in the intimal layer (75) and adhesion of leukocytes to the endothelium (3).

Systemic inflammation is a well-established feature of OSA (144, 345, 412, 564, 609, 699, 749). Increases in some OSA-related markers of inflammation, especially tumor necrosis factor-α (TNF-α), are thought to be caused by sleep disruption because they are correlated with daytime sleepiness (564, 699, 700). Nevertheless, the primary proatherogenic feature of OSA appears to be intermittent hypoxia. In a large group of OSA patients without known cardiovascular disease, nocturnal oxygen saturation levels were predictive of carotid artery thickening and plaque occurrence, independently of hypertension (27). In another study, desaturation index (not AHI) was the strongest predictor of plasma levels TNF-α and interleukin-8 in patients with OSA (565).

In addition to its role as an initiator of atherogenesis, OSA may contribute to progression of established lesions. Multiple sources of evidence suggest that OSA promotes thrombosis, because enhanced platelet activation and aggregation (60, 149, 411, 559, 578, 702), enhanced erythrocyte adhesiveness and aggregation (497), increased fibrinogen levels (103, 145, 497, 646, 717), and diminished fibrinolytic activity (538) have all been observed in patients with OSA. Increased endothelial cell apoptosis in patients with OSA may also contribute to coagulation abnormalities (150). OSA-induced inflammatory chemokines and cytokines are potential contributors to plaque rupture via their effects on extracellular matrix (75). The hemodynamic perturbations caused by repetitive apneas may destabilize vulnerable plaques and may enhance oscillatory shear stress and superoxide production (225) at areas of the vascular tree predisposed to atherosclerosis.

8. Cerebrovascular disease

An association exists between OSA and atrial fibrillation (184, 212, 291); therefore, thromboembolism may be an important cause of stroke in OSA patients. Moreover, Beelke et al. (44) recently demonstrated that apnea-induced changes in intrathoracic pressure cause intrathoracic shunting in patients with patent foramen ovale. In the setting of hypercoagulability, this anomaly could give rise to embolization. Recent case reports implicate OSA as a possible cause of cryptogenic stroke (476).

Individual episodes of OSA cause marked fluctuations in cerebral blood flow (31, 217, 309) (see above). Diminished cerebrovascular reactivity to hypoxia and hypcapnia has been observed in patients with OSA (133, 175, 545). Similar impairments are seen in patients with ischemic cerebrovascular disease (111, 383); however, whether these alterations in vascular regulation contribute to the pathogenesis of stroke in the setting of OSA is unknown.
9. Alterations in cardiac function and structure

Episodes of OSA activate hypoxia-related neurohumoral pathways, increase transmural pressures and afterloads, and lead to chronically elevated blood pressure, any of which could contribute to the development or worsening of ventricular dysfunction. In addition, fixed increases in large artery stiffness (139, 510) may lead to cardiac remodeling. Increased left ventricular mass and diastolic dysfunction have been observed in patients with moderate to severe OSA (140, 179, 232, 475), but not in those with less frequent apneas and hypopneas (448). In some of these studies, left atrial size was increased as well, which could explain the association between OSA and atrial fibrillation (184, 484).

What is the trigger for OSA-related ventricular dysfunction? Negative intrathoracic pressures resulting from OSA episodes, via their effects on left and right ventricular afterloads, may contribute to ventricular remodeling. Nevertheless, animal models indicate that chronic exposure to intermittent hypoxia per se, in the absence of airway obstruction, is also sufficient to impair ventricular function (95, 97, 230). In these models, ventricular dysfunction and remodeling were accompanied by markers of oxidative stress (95, 230). On the cellular level, both hypertrophy and apoptosis of cardiac myocytes have been observed following CIH in rats (97).

10. Genetic aspects of OSA-related cardiovascular disease

The fact that cardiovascular disease is not a universal finding in patients with OSA suggests a role for genetic predisposition. On the basis of what is known about the mechanisms of cardiovascular disease in OSA, several genetic candidate genes have been investigated.

Several investigators have studied the role of angiotensin converting enzyme (ACE) gene polymorphisms (61, 490, 766). In aggregate, these studies suggest complex, potentially important interactions between ACE gene insertion/deletion polymorphisms and SDB as mechanisms for OSA-related hypertension. The ACE D allele is associated with hypertension in subjects with mild-moderate OSA, whereas in patients with severe OSA, the ACE D allele may have a protective influence (61, 490).

An association between OSA and a leptin receptor gene polymorphism has recently been reported (524). In this study, total cholesterol, high-density lipoprotein (HDL), low-density lipoprotein (LDL), and triglyceride levels were higher in OSA patients with the Arg/Arg genotype. OSA is also associated with TNF-α and β2-adrenergic receptor polymorphisms (38, 555).

Polymorphism in the haptoglobin gene, which is associated with cardiovascular risk in diabetes (351), was recently recognized as a risk factor for OSA-related cardiovascular disease (344). Putative mechanisms for this association include impairments in the immunomodulatory and antioxidant properties of haptoglobin (344). Another potentially fruitful target of investigation is the extracellular superoxide dismutase gene. A common variant of this gene is associated with increased risk of coronary disease in non-SDB populations (287); however, the relationship between this gene variant and cardiovascular disease in OSA has not, to our knowledge, been explored. Clearly, further studies are required to establish the genetic variations and the gene-environment interactions responsible for increasing the risk of cardiovascular disease in patients with SDB.

E. Summary: Cardiovascular Sequelae of Sleep Apnea

A schematic representation of the current concept of interdependent mechanisms by which OSA in humans and CIH in animal models led to cardiovascular dysfunction and disease is shown in Figure 15. In the carotid body and in brain regions influencing sympathetic outflow, intermittent hypoxia causes upregulation of ANG II/NADPH oxidase and downregulation of NOS. The resultant excess of superoxide ion results in chronically elevated sympathetic outflow. Sympathetic overactivity, in turn, produces trophic and pro-atherosclerotic effects on resistance vessels via oxidative stress and inflammation. Increased sympathetic outflow to the kidney stimulates renin release and leads to elevated circulating levels of ANG II and aldosterone, two hormones with oxidative stress and inflammatory effects of their own. Sympathetic activation of liver and spleen, T-cell-rich tissues, may play a role in the inflammatory response. In OSA, episodic hypercapnia and arousal from sleep play secondary roles by potentiating the hypoxia-induced increase in sympathetic outflow (248, 422, 423).

In the years since identification of the syndrome, much has been learned about the cardiovascular sequelae of OSA; nevertheless, many questions remain unanswered. Do these sequelae have functional, as well as disease-related, consequences (e.g., do they impair regulation of the systemic or pulmonary circulation during exercise)? Can OSA patients with increased risk for development of cardiovascular disease be identified on the basis of genotype or phenotype, thereby enabling rational decisions about who to treat? Can the cardiovascular consequences of OSA be prevented or ameliorated by pharmacological interventions (e.g., antioxidant or anti-inflammatory agents)?
VI. OBSTRUCTIVE SLEEP APNEA AND INSULIN RESISTANCE

A. Introduction

Insulin resistance is a central part of the metabolic syndrome, a condition that is reaching epidemic proportions in Western Society and now emerging in developing countries (298, 530). The metabolic syndrome has many features in common with OSA including obesity, hyperlipidemia, hypertension, and insulin resistance. OSA is so interwoven in the fabric of the metabolic syndrome, or Syndrome X, that the combination of OSA and metabolic syndrome has been labeled “Syndrome Z” (723). Consequently, the causal nature of the relationship between OSA and various components of the metabolic syndrome, in particular insulin resistance, has been difficult to untangle.

As detailed above, there is now a considerable body of evidence indicating that OSA can independently contribute to the development of sustained daytime hypertension. In contrast, the concept that OSA potentially impairs insulin sensitivity (i.e., causes insulin resistance), is a much more recent development. Earliest reports that OSA may impair glucose homeostasis, independent of obesity, began to surface in the early 1990s, and the field has slowly gained momentum since. However, the lack of preclinical data to guide and support the clinical studies, in addition to the technical difficulties associated with assessing metabolic end points, has hampered progress in the field of OSA and insulin resistance.

The measurement of insulin sensitivity, or surrogates of insulin sensitivity, require invasive measurements that are often difficult and technically challenging. Even the simplest marker of insulin sensitivity, the homeostasis model assessment (HOMA) index, requires measurement of blood glucose and plasma insulin. Other metabolic assessments utilized in OSA patients include the standard clinical assessment of oral glucose tolerance test (OGTT) and the predominantly research-focused, frequently sampled intravenous glucose tolerance test (IVGTT). The latter test provides several parameters of insulin and glucose homeostasis, including a modeled estimate of insulin sensitivity (51). Due to technical demands, very few studies have utilized the gold standard measurement of insulin sensitivity: the hyperinsulinemic euglycemic clamp (125). Thus the challenges of measuring insulin sensitivity and other metabolic parameters involved in glucose homeostasis have acted to impede research in this relatively new field.

B. Prevalence and Incidence of Insulin Resistance Type 2 Diabetes in OSA

The first studies examining the prevalence of glucose dysregulation and OSA appeared in 1993. Since this time, a group of studies, generally involving large epidemiological cohorts, have used questionnaire-based surrogates of OSA, such as snoring or witnessed apnea, and assessed the prevalence or incidence of diabetes based on elevated glucose levels, medication use, or self-reported diabetes. The largest of these studies was a prospective cohort from the Nurses’ Health Study (4) that followed 69,852 women without diagnosed diabetes over a 10-yr period. During the course of the 10-yr follow-up period, 1,957 women were diagnosed with the development of type 2 diabetes. The presence of snoring was associated with a 2.25 increase in relative risk of developing diabetes compared with nonsnorers, and the association remained significant after adjustment for covariates including BMI, activity, smoking, and family history of diabetes. These results indicated that in a large population-based study
snoring is independently associated with an elevated risk of developing type 2 diabetes (4).

Since snoring is only a surrogate marker of OSA, other epidemiological studies have used polysomnography to objectively classify the presence and severity of OSA. In general, these studies have involved smaller cohorts recruited from clinic populations. Two of the larger studies with clinic-based sample sizes of 250–300 subjects both demonstrated a positive association, independent of obesity, between the severity of OSA and indexes of insulin resistance determined by fasting insulin and glucose (268, 652). Similar findings were reported by Punjabi et al. (535) from a community-based sample of overweight or obese, but otherwise asymptomatic, males. In this study by Punjabi et al. (535), subjects with mild or moderate to severe OSA had significantly increased odds ratios for elevated fasting and 2-h glucose levels from the OGTT, after adjustment for both BMI and percent body fat. A large community-based pediatric study of 907 children also found a significant association between an AHI of >5 and elevated circulating insulin and HOMA index of insulin resistance (542). By far the largest epidemiological study to date that directly assessed OSA by polysomnography and measured glucose and insulin levels under fasting conditions and after an OGTT was based on a subset of 2,656 subjects from the ongoing Sleep-Heart-Health study (534). In subjects with an AHI of 15 events/h or greater, there were small, but statistically significant, elevated odds ratios of 1.46 for fasting glucose and 1.44 for 2-h glucose levels in the OGTT, after adjustment for age, BMI, waist girth, race, sex, and smoking. In addition, the HOMA index of insulin sensitivity was also significantly elevated in subjects exhibiting an AHI of 15 events/h or greater. In combination, the prevalence data from multiple epidemiological studies support an independent association between OSA and impaired glucose homeostasis.

Despite several positive prevalence studies of OSA and indexes of insulin resistance, there is so far only one prospective study examining the association between OSA, determined by polysomnography, and the development of type 2 diabetes (546). Comparable to previous cross-sectional studies, they showed a positive association between clinically significant OSA and a diagnosis of type 2 diabetes in 1,387 participants in the Wisconsin Sleep Cohort after adjustment for age, sex, and waist girth. However, in a follow-up study of 978 subjects, the odds ratio for developing type 2 diabetes within a 4-yr period for those with an AHI of >15 events/h did not reach statistical significance after adjustment for waist girth. Although many factors may account for the potential disparity between the cross-sectional and longitudinal findings of this study, the results suggest that epidemiologically there is a lack of strong support for a causal relationship between OSA and the development of insulin resistance and type 2 diabetes. An alternative approach using prospective studies in clinical populations to examine the existence of causality is to determine the impact of therapeutic treatment of OSA on glucose and insulin regulation.

C. Effect of Treatment of OSA on Insulin Sensitivity

Nasal continuous positive airway pressure (nCPAP) is the predominant therapeutic treatment for OSA and is highly effective at eliminating periods of airway obstruction during sleep. The relationship between the use of nCPAP and indices of insulin resistance has been a focus of several recent studies in OSA patients. The studies have been predominantly of small sample size with subjects recruited from sleep or diabetes clinics; the length of treatment as variable as 1 day to 6 mo; the nCPAP intervention uncontrolled with limited adherence data; and few studies utilized the hyperinsulinemic euglycemic clamp as the major outcome variable. In general, the studies were largely negative with variable lengths of nCPAP treatment having no effect on fasting insulin levels (119), glucose tolerance (104), or insulin sensitivity (629). One exception was the 4-mo nCPAP trial by Brooks et al. (71), which did show a 32% increase in insulin sensitivity using the hyperinsulinemic euglycemic clamp in subjects with effective nCPAP treatment. Thus this initial small and disparate group of treatment studies did not appear to support the relatively larger body of prevalence studies, suggesting an independent effect of OSA on disrupting glucose homeostasis.

However, a more recent and larger study by Harsch et al. in 2004 (226) demonstrated a positive effect of nCPAP on insulin sensitivity. In an uncontrolled longitudinal study in 40 OSA patients, insulin sensitivity improved 18% after just 2 days of treatment and by 31% after 3 mo of treatment. Interestingly, a post hoc analysis showed that subjects with a BMI of under 30 kg/m² showed much greater improvements in insulin sensitivity within 2 days of treatment compared with subjects with a BMI at or above 30 kg/m². In a follow-up study in nine of the subjects who were adherent to nCPAP, the improvement in insulin sensitivity was still evident on average 2.9 yr later (586). Thus the concept has emerged that the detrimental effects of OSA on insulin sensitivity are potentially more apparent in the absence of comorbidities associated with obesity. Interestingly, in a prepubertal pediatric population, the reverse scenario was recently reported in response to a therapeutic intervention. Circulating insulin and the insulin-glucose ratio were significantly decreased 6–12 mo following adenotonsillectomy in obese subjects, but not nonobese subjects (202). The interaction between the effects of therapy and comorbid-
ties on insulin resistance in OSA will require careful consideration in future studies.

A randomized, placebo-controlled nCPAP study by West et al. (719) has recently challenged the positive findings of Harsch et al. (226). Three months of nCPAP therapy in patients with known type 2 diabetes and newly diagnosed OSA showed an improvement in measures of sleepiness, but there was no change in HbA1c, HOMA index, or insulin sensitivity as measured by the clamp procedure in either the therapeutic or placebo nCPAP groups. A difference between these two conflicting studies is that West et al. (719) used more obese subjects with preexisting type 2 diabetes. However, a similar 6-wk randomized, placebo-controlled nCPAP study in nondiabetic patients reported no improvements in metabolic outcomes with therapy, although there was a trend ($P = 0.08$) for HOMA to improve by ~15% (109). Both nCPAP studies reported average compliance rates between 3–4 h per night, which may be critical given the report by Dorkova et al. (136) demonstrating significant improvements in HOMA-assessed insulin resistance in nCPAP compliant (average 5.07 h/night) but not noncompliant (average 3.49 h/night) subjects after 8 wk. It remains to be seen whether an appropriately powered, placebo-controlled, nCPAP study in OSA patients without overt diabetes and long-term follow-up can show significant benefits of therapy on insulin resistance. In summary, the clinical evidence for a causal pathway between OSA and insulin resistance remains equivocal. Although prevalence data for an association between OSA and insulin resistance exists, incidence studies and interventional therapeutic studies have not consistently supported a role for OSA inducing insulin resistance.

D. Experimental Evidence That OSA Can Lead to Insulin Resistance

Two primary physiological disturbances that characterize OSA, acute periods of hypoxic stress and disruption of sleep, can potentially impair glucose homeostasis. Several epidemiological studies including the Sleep-Heart-Health Study (534) have reported that fasting basal glucose levels, the HOMA index, or blood glucose levels during the OGTT are elevated as a function of the degree of nighttime hypoxemia in OSA. Exposure to chronic hypoxia, as occurs with ascent to altitude, can, at least initially over the first 48 h, lead to insulin resistance (338). However, by 7 days of exposure to 4,559 m above sea level, insulin sensitivity is correcting back to sea level values. Experimentally, acute insulin resistance, as determined by a euglycemic clamp, reached a maximum 20 min after a continuous 30-min exposure to hypoxia that lowered arterial oxygen saturation to ~75% in normal individuals (Fig. 16) (463). Thus short-term exposure to sustained hypoxia produces insulin resistance in humans, but the acute, repetitive hypoxic stress that occurs during sleep in OSA may have different metabolic consequences. The potential specificity of the paradigm of hypoxic stress associated with OSA has prompted studies of glucose homeostasis and insulin resistance in rodent models of IH. Interestingly, it appears that glucose homeostasis in mice is a time-dependent phenomena affected by the presence or absence of the IH stimulus (748). Blood glucose levels and insulin resistance as assessed by the hyperinsulinemic euglycemic clamp are elevated during

![Fig. 16. Hyperinsulinemic euglycemic clamps performed in healthy humans and mice during exposure to hypoxia.](http://physrev.physiology.org/)
exposure to IH during the light or sleeping phase (Fig. 16) (263). In contrast, during the 12-h dark or active period in which animals are maintained in a nonhypoxic constant room air environment, blood glucose and insulin resistance may actually be decreased below control levels (522), suggesting an overcompensation in insulin sensitivity. Thus the degree of insulin resistance may fluctuate on a diurnal basis dependent on the presence or absence of the IH stress. Diurnal fluctuations in insulin resistance represent a very different pattern of response compared with the development of hypertension in rodent models of IH, where arterial blood pressure remains continuously elevated (329, 664). Thus the mechanisms by which IH chronically elevates blood pressure, but phasically impacts on insulin resistance, suggest differing downstream mechanisms produce the cardiovascular and metabolic disruptions.

Since OSA and obesity frequently coexist, the disruptive effects of IH stress on glucose homeostasis may be potentially exacerbated by adiposity. The comorbid impact of obesity was examined in a rodent model of IH. Genetically obese ob/ob mice exhibited increasing basal insulin levels associated with impaired glucose tolerance over a 3-mo period of exposure to IH compared with weight-matched control ob/ob mice (522). The confounding or interacting effects of obesity have likely contributed to much of the inconsistency in the clinical literature where comorbidity associated with obesity is often present.

Recently, techniques have been developed to assess the impact of experimental IH on glucose homeostasis in healthy humans free of OSA and metabolic dysfunction. The study by Louis et al. (372) showed that a 5-h period of experimental IH in normal sleeping humans (20–30 hypoxic events/h) decreased insulin sensitivity, as assessed by the IVGTT, from 3.8 to 2.6 mU·l⁻¹·min⁻¹. These data, in combination with the rodent studies, suggest that acute exposure to IH can cause insulin resistance in both humans and animals even in the absence of comorbid conditions.

In addition to hypoxic stress, impaired sleep is also a potential candidate for metabolic dysfunction. Reducing sleep time to 4 h/night over a 6-day period decreases the rate of glucose clearance, glucose effectiveness, as well as the acute insulin response to glucose (639). There are also several prospective epidemiological studies, including the large Nurses Health Study mentioned above (19), demonstrating that short sleep duration increases the risk of developing diabetes. Furthermore, sleep deprivation can reduce leptin levels and increase ghrelin levels, potentially acting to stimulate appetite (640, 665). Thus the impact of sleep deprivation on compromising metabolic function may be exacerbated by a secondary effect to increase appetite, which itself may lead to weight gain and further metabolic dysfunction.

Sleep deprivation, or sleep restriction, may not reflect the disturbances in sleep that occur in OSA. Typically, total sleep time is not significantly restricted in OSA, but rather sleep is fragmented by repetitive arousals resulting from the impaired breathing during sleep. The study by Stamatakis et al. (643) attempted to model the sleep fragmentation of OSA by arousing normal healthy humans from sleep over two nights (30–40 times/h) using auditory and mechanical stimuli. This form of experimentally induced sleep fragmentation caused a 20.4% decrease in the insulin sensitivity index as assessed by the IVGTT. There does not appear to be any comparable animal models of sleep fragmentation that have demonstrated metabolic abnormalities. Clearly there is a need for more clinical and translational research to determine whether sleep fragmentation can contribute to the development of insulin resistance.

E. Are There Plausible Mechanisms for OSA to Cause Insulin Resistance?

Determining the mechanisms that cause insulin resistance is a focus for a large number of researchers in the area of type 2 diabetes, although currently few approach the issue from the perspective of OSA. For the purposes of this review, potential mechanistic pathways of insulin resistance are split into two broad categories defined as “classical” and “lipotoxic,” respectively, in Figure 17. These categories and pathways should be considered neither exhaustive nor mutually exclusive, but rather provide a framework for exploring the mechanisms through which OSA can putatively cause insulin resistance. In the absence of obesity, several factors listed under the classical pathways that lead to insulin resistance are relevant to OSA.

The ability of OSA to activate the sympathetic nervous system is now well characterized. Increased sympathetic nerve activity is implicated as a primary mechanism in the development of sustained hypertension in OSA patients (83, 634) and rodent models of IH (170). Since activating the sympathetic nervous system can also potentially impact on insulin sensitivity (246), it has been proposed that increased sympathetic nerve activity may lead to insulin resistance in OSA patients (226). Although no clinical data exist to support or refute this hypothesis, a study in conscious mice demonstrated that the development of insulin resistance over a 9-h period of IH exposure persisted even after complete pharmacological denervation of both the sympathetic and parasympathetic nervous systems with a ganglionic blocking agent (263). This animal study suggests, at least in the short term, that activation of the sympathetic nervous system is not required for the development of insulin resistance in response to hypoxic stress. However, the possibility remains that hyp-
oxic activation of the sympathetic nervous system, or an increase in circulating catecholamines, contributes to the long-term progression of insulin resistance and metabolic function that may occur over decades in patients exhibiting OSA and obesity.

Increased circulating catecholamines are just one of a group of circulating hormones that act in a counterregulatory fashion to the blood glucose-lowering actions of insulin (246), as well as inhibit insulin release from the pancreas (525). Clinical studies in OSA patients (172, 408) and rodent studies of IH (348) have demonstrated increased levels of circulating catecholamines. However, no study has attempted to link an increase in catecholamines to changes in insulin sensitivity or glucose homeostasis. Growth hormone is another counterregulatory hormone that is affected by sleep and OSA. In contrast to catecholamines, growth hormone levels appear, if anything, to be reduced in the presence of OSA (191, 569), suggesting that this hormone does not play any direct role in the development of insulin resistance. However, growth hormone is the predominant factor controlling release from the liver of insulin-like growth factor I (IGF-I), a peptide with insulin-sensitizing actions. Studies in adult and pediatric patients show that OSA decreases IGF-I and that treatment with nCPAP in adults (211) or surgical adenotonsillectomy in children (34) can increase IGF-I. In contrast to these positive therapeutic effects, a more recent study in adult male OSA patients was unable to detect an independent effect of treatment to increase IGF-I in a parallel, randomized, sham placebo-controlled 1-mo nCPAP trial (403). Furthermore, the relationship between any OSA-induced lowering of IGF-I and the development of insulin resistance has not been directly explored.

Intuitively, the hypoxic stress of OSA would likely activate the hypothalamic-pituitary-adrenal (HPA) axis, elevate cortisol levels, and putatively contribute to insulin resistance. Rodent studies of IH have demonstrated sensitization of the HPA axis (377) as well as increased levels of corticosterone, the predominant glucocorticoid in rodents, that peak during the 12 h of the light or sleeping period when the hypoxic stimulus is present (748). Moreover, these spikes in corticosterone mirrored spikes in glucose, an association that suggests a potential contribution to hypoxic-mediated insulin resistance. However, there are few human studies that have examined cortisol changes in OSA. What few studies have been conducted indicate that OSA does not affect cortisol levels (156, 211, 403). However, one study has shown an increase in cortisol levels in OSA patients relative to weight-matched controls, and the elevated cortisol levels in patients were reduced with nCPAP (68). Interestingly, the study by Speigel et al. (639) in normal healthy young adults demonstrated that sleep restriction to 4 h/night changed the circadian profile of circulating cortisol with elevated levels in the afternoon and early evening. Thus similar to the somatotropic axis, it may be necessary to assess any impact of OSA on the HPA axis hormonal profiles across the entire circadian cycle.

The development of insulin resistance and type 2 diabetes is largely dependent on the presence of obesity. There is now a growing body of evidence that the “lipotoxic” effects of obesity play an important role in the pathogenesis of insulin resistance. Supporting this hypothesis are observations that acute hyperlipidemia induces insulin resistance (20, 59) and that decreasing the metabolic availability of lipids in vivo increases insulin sensitivity (457, 661). Proinflammatory/stress pathways have been proposed as an important link between lipotoxicity and the development of insulin resistance (Fig. 17). Cellular responses to inflammatory and stress signals are mediated by a number of ubiquitously expressed signaling cascades, including the NF-κB pathway. Increased activity in proinflammatory/stress pathways has been implicated in impairing insulin action in peripheral tissues (762).

Multiple factors have been proposed to activate proinflammatory/stress pathways in obesity, including generation of reactive oxygen species, release of inflammatory cytokines, hyperlipidemia, and ectopic deposition of fat. Interestingly, all of these pathways are potentially activated by the hypoxic stress of OSA in patients or experimentally induced IH in rodents. The hypoxic stress of OSA is a unique stimulus that incorporates a rapid period of tissue deoxygenation immediately followed by rapid tissue reoxygenation. These rapid and repetitive swings in deoxygenation/reoxygenation have the poten-
tial to generate reactive oxygen species (144, 594) and lead to lipid peroxidation (410) in OSA patients and in multiple organs including liver (357), heart (95), and brain (694) in rodents exposed to IH. In addition to any direct actions of hypoxic stress from OSA, lipotoxicity may also generate reactive oxygen species and ultimately activate proinflammatory/stress pathways that lead to insulin resistance. Now evidence is emerging that OSA produces a proinflammatory state, and specifically that OSA and IH can activate the NF-κB pathway (205, 564, 565).

Adipocytes are a major source of circulating cytokines that both induce and respond to proinflammatory stress pathways. In general, cytokines are secreted into the circulation as a function of the size of an adipocyte, consequently establishing a positive relationship between adiposity and circulating cytokines (29, 623). For example, two important inflammatory cytokines, TNF-α and IL-6, have elevated circulating levels in obesity and are decreased by weight loss (416). Clinical studies suggest that OSA may have an independent role in further increasing circulating levels of TNF-α, IL-6, as well as the general inflammatory marker C-reactive protein, above the levels seen in obese, nonapneic control subjects, as well as in OSA patients treated with nCPAP (409, 699, 749). Apart from leptin (522), an insulin-sensitizing cytokine, there are little data assessing whether hypoxic stress increases circulating levels of inflammatory cytokines in rodent models of IH.

Both clinical studies and animal studies suggest that an independent relationship may exist between OSA and hyperlipidemia. In a sample of nearly 5,000 subjects from the Sleep Heart Health study, there was a positive association between the severity of OSA and increased serum total cholesterol and triglycerides, as well as decreased serum HDL, in men and women aged less than 65 yr (443). Other smaller clinical studies involving nCPAP support a role for OSA increasing HDL and lowering LDL cholesterol (102, 559). Furthermore, hyperlipidemia can result from exposure to IH in rodent models (358). Thus the hypoxic stress of OSA potentially increases the risk of hyperlipidemia.

In addition to increasing circulating lipid levels, there is evidence that hypoxic stress may affect the accumulation of lipids and subsequent inflammation in organs, as well as the overall distribution of body fat. Exposure of mice to IH can lead to lipid accumulation in the liver (358), upregulation of transcription factors controlling lipid biosynthesis in the liver (355), cause lipid peroxidation and activation of the NF-κB pathway (355), and cause steatohepatitis in a mouse model of diet-induced fatty liver (584). Clinical studies also suggest that OSA is a potential risk factor for steatosis, elevated liver enzymes, and steatohepatitis (668). In addition to producing ectopic fat accumulation and subsequent inflammation in the liver, there is indirect evidence that OSA may predispose to weight gain (506) and potentially influence the distribution of visceral versus subcutaneous fat accumulation (104). Although no studies have focused on whether OSA or IH can lead to ectopic fat accumulation in muscle, the predominant source of insulin-mediated glucose uptake, current evidence suggests that lipid accumulation can occur in the liver and is associated with a proinflammatory state.

In summary, the downstream sequelae of OSA impact a vast array of organ systems and cellular processes in ways that could lead to the development of insulin resistance. The question becomes not “are there plausible mechanisms for OSA to cause insulin resistance?” but rather “what is the relative importance of the many mechanistic pathways through which OSA may cause insulin resistance?” Despite the challenges imposed by tackling this question, there remain many other issues that hinder the development of this area of research.

F. Future Challenges

There is an immediate need for large-scale incidence studies of insulin resistance and type 2 diabetes in well-characterized patients with polysomnographic determination of OSA. The more sophisticated assessment of insulin resistance (e.g., hyperinsulinemic euglycemic clamp or IVGTT), the more significance such studies will carry. Similarly, long-term correctly controlled nCPAP studies are required to assess the relative burden of disease and determine the potential therapeutic benefits associated with treatment, as well as address the fundamental question of whether insulin resistance and type 2 diabetes are reversible by a therapy that is highly effective at eliminating respiratory disturbances during sleep. Moreover, are there subgroups of pediatric and adult patients, such as those with preexisting type 2 diabetes or increased visceral fat accumulation, who are more resistant to the therapeutic metabolic benefits of nCPAP? The effectiveness of nCPAP for improving insulin sensitivity should be compared quantitatively with the insulin-sensitizing effects of pharmacological therapies and behavioral therapies of weight loss and physical activity. Study designs need to become more comprehensive with respect to multiple sampling across the day and night, as well as controlling potentially confounding factors such as food intake and physical activity. However, despite all these challenges, the issue that restricts progress the most is the overriding and pervasive influence of obesity in both metabolic disorders and OSA. Solving the algebraic equation “Z – X” and extracting the Z component from Syndrome X is proving extremely difficult. Whereas animal models can provide unique insights and control for the effects of obesity, ultimately discoveries need to be translated back into the clinical arena. The data gathered to date suggest the potential for disturbances in sleep and breathing that occur in
OSA to possess some degree of independent action, but any action likely represents the tip of a metabolic iceberg buoyed by obesity and insulin resistance.

VII. NEURAL INJURY IN OBSTRUCTIVE SLEEP APNEA

A. Introduction

The majority of adults with untreated OSA present to the clinician with one or more neurobehavioral impairments, including sleepiness, fatigue, depressed mood, impaired memory, and/or poor concentration. Less commonly, individuals presenting with OSA note motor and/or sensory impairments. It has been difficult to discern in clinical trials whether OSA directly contributes to any one of the associated neurological complaints. This is, in part, because many individuals with OSA have comorbidities that are associated with neural injury, including diabetes, hypertension, and cerebrovascular disease, as described in the previous sections. Moreover, the onset of sleep apnea is insidious, and many individuals present to the physician for evaluation of sleep apnea only years after symptoms were first noted. Nonetheless, the concept that OSA contributes to neural injury is strongly supported by the observation that effective treatment of OSA with continuous airway pressure frequently improves many of the neurological signs and symptoms. Severe neurobehavioral impairments, however, typically do not fully reverse. This raises an important clinical question: does OSA result in irreversible neurobehavioral sequelae?

B. Insight From Neuroimaging Studies

Neuroradiography can provide important insight into structural and functional differences associated with disease in a noninvasive manner. In the case of obstructive sleep apnea, where CPAP affords effective treatment of the disease, neuroimaging before and after treatment may also provide insight into direct effects (reversible effects) of the disease on the CNS. Before reviewing the research findings of neuroimaging in sleep apnea, it is important to consider the difficulties and limitations of such studies. One of the challenges in interpreting findings in neuroradiological studies concerning the effects of sleep apnea on brain structure or function is that comorbid conditions (e.g., cardiovascular disease, diabetes) also negatively impact on brain function. For example, there is no general consensus whether subjects should be matched for all other diseases so that the only difference across groups is a high or low apnea index, or whether studies should simply compare all subjects, regardless of comorbidities, with high versus low apnea hypopnea indexes. As you will see below, the two strategies can lead to very different interpretations of the impact of sleep apnea on brain function. A second challenge is an inherent challenge with associative and cross-sectional studies: whether the neural injury preceded sleep apnea or vice versa. A critical obstacle is that there may well be interindividual differences with susceptibility to neural injury such that not all subjects with sleep apnea will have significant neural injury. Presently for sleep apnea we cannot define these vulnerable subsets. Until the groups of individuals at risk are defined, very large sample sizes will be required to detect overall differences; unfortunately, to date, most imaging studies involve sample sizes smaller than 30. With these limitations in mind, there have been several important, insightful studies.

Three groups of researchers have examined grey matter loss in sleep apnea. Macey and co-workers (379, 380) used MRI to examine grey matter in 21 individuals with sleep apnea and 21 controls (all male, ranging from 28–70 yr of age). They found significant reductions in grey matter in several brain regions, including the hippocampus and cingulate cortex. More importantly, grey matter loss correlated positively with apnea severity. Morrell et al. (429) also found grey matter loss in the hippocampus in OSA. In contrast, O’Donoghue et al. (456) examined 27 males with severe OSA and 24 male controls and did not find differences in grey matter. In this study, subjects were matched for diabetes, hypertension, and other cardiovascular diseases. It is possible, as shown below in animal models, that mechanisms involved in cardiovascular and diabetic injury in sleep apnea are the same as brain injury. Thus O’Donoghue et al. (456) may have selected out the individuals who are most predisposed to neural injury. An alternative explanation for the negative results is that the O’Donoghue study recruited younger adults and thus may have missed effects that require a longer course of sleep apnea or an older age for measurable injury.

Positron emission tomography (PET) of the brain can provide insight into regional brain metabolism and may also provide insight into abnormalities in specific neurochemical transmission. While the latter has yet to be taken advantage of in sleep apnea studies, there is a recent report highlighting the value of PET scanning in sleep apnea (14). In this study, subjects with unexplained residual sleepiness despite effective therapy for sleep apnea with CPAP were examined for fluoro-deoxyglucose uptake in the prefrontal cortex in parallel with polysomnography and a vigilance test. Four of the seven subjects had impaired glucose utilization in the frontal and/or temporal cortex; one additional subject had reduced glucose utilization in the parietal cortex. The remaining two subjects had no obvious PET scan abnormalities. Of clinical significance, this study supports the concept that some
cognitive impairments, including sleepiness, are not fully reversible in all patients with sleep apnea.

Vascular injury and disturbances in blood flow may contribute to neural loss. In a recent study by Minoguchi et al. (411), brain MRI was used to compare the percentage of silent brain infarctions in subjects with and without OSA. Remarkably, 25% of individuals with severe sleep apnea had infarctions. In contrast, only 7% of obese matched nonapneics had infarcts evident. In support of a direct relationship, the group measured two serum markers for cerebrovascular disease, sCD40L and sP-selectin. Both of these markers were high in sleep apnea and declined upon effective treatment with nasal CPAP. Recently, single photon emission tomography (SPECT) was implemented to measure regional blood flow in severe sleep apnea (284). In this study 27 subjects with severe OSA (AHI, 30 – 104/h) were compared with 27 controls, age and sex matched. As with lesions, blood flow appears regionally modified in awake subjects with OSA, with lowest levels in the parahippocampal gyri, pericentral gyri and cuneus. Whether this impacts cognitive function during wakefulness or is secondary to neural injury is not clear.

MRTI with spectroscopy may be used to examine metabolic disturbances, where disturbances include neuronal loss, gliosis, and other causes of altered metabolism. A reduced N-acetylaspartate/NAA-to-choline ratio was found for the cerebral white matter, where the AHI correlated negatively with the NAA-to-choline ratio (290).

These studies prompt the question: Is this loss of brain tissue associated with impaired cognitive performance? Several recent studies have begun to address this issue. Thomas et al. (675) examined cognitive function in parallel with functional MRI in individuals with severe OSA. Ages of the enrolled subjects ranged from 21 to 50 yr, with a male predominance. Subjects with OSA had less activation of the prefrontal cortex while performing a working memory task. A similar decrement was observed in hypoxic and nonhypoxic subjects with sleep apnea, suggesting that hypoxia does not influence the decrement in prefrontal cortical activation during learning. In contrast, hypoxic subjects showed far less activation in the parietal cortex. This suggests that within the brain there are regional differences for hypoxia-sensitive neuronal tissue and arousal or sleep disruption-sensitive neuronal tissue. One of the more alarming findings from recent studies is that young children with sleep apnea may also have neuronal loss and cognitive impairments. Hallowes et al. (218) examined 19 children with sleep apnea and 12 controls, matched for age, gender, ethnicity, and socioeconomic class. Children with severe sleep apnea had significant decrements in their IQ (15 points) and significant decrements in verbal working memory and verbal fluency. These cognitive impairments paralleled reductions in the NAA-to-choline ratio in the hippocampus and frontal cortex by spectroscopy. These findings have serious clinical implications and are worthy of confirmation and then determination of reversibility. At the very least, these findings should prompt careful screening for sleep apnea in all obese children and an aggressive campaign to educate parents on the risks of childhood obesity and to increase efforts to reduce childhood obesity. If these findings prove irreversible, the sleep community should play an important role in campaigning these issues.

C. Evidence of Neural Injury From Animal Models

While there are numerous physiological perturbances in OSA that might disturb neuronal homeostasis and function, David Gozal was one of the first researchers to explore whether the frequent hypoxia/reoxygenation patterns in sleep apnea result in lasting neural injury and impaired neural function. To test this, Gozal et al. (203) exposed young adult rats to either 2 wk of fluctuating ambient oxygen patterns modeling oxygenation in severe sleep apnea, constant hypoxia of the same duration, or room air oxygen tension and then examined the effects of varied oxygenation on learning and neuronal health (203). Gozal et al. (203) identified learning impairments and increased apoptosis within the CA1 region of the hippocampus only in rats exposed to intermittent hypoxia. Similarly, spatial memory impairment was observed several months after intermittent hypoxia exposure to newborn rat pups (123, 124). Thus memory impairments may occur as a result of hypoxia/reoxygenation patterns modeling OSA, and these may persist well beyond the hypoxia/reoxygenation exposure. Future clinical trials should now focus on spatial memory function in individuals with OSA and oxyhemoglobin desaturations. At the same time, animal studies should identify the mechanisms by which frequent hypoxia/reoxygenation events result in hippocampal injury and dysfunction.

D. Daytime Sleepiness

One of the most common neurobehavioral impairments in OSA is sleepiness. In fact, two-thirds of adults with OSA complain of significant sleepiness and/or fatigue (100). When treated for OSA, patients typically report less somnolence (153, 491). Despite marked improvements in subjective sleepiness, randomized controlled trials show small improvements in objective sleepiness (<1 min increase in mean sleep latency overall) despite effective therapy for OSA (491). This objective measure is the average latency to fall asleep during four or five nap opportunities distributed across the morning and afternoon. Thus falling asleep 1 min later across four or five nap opportunities is not a clinically significant improvement in wake function. Although apneic events result in many physiological disturbances as reviewed in previous sections, the oxygen desaturation indexes most strongly
predict sleepiness, relative to other polysomnographic parameters, including sleep time, AHI, or arousal index (42, 114, 153, 303, 678). While clinical studies show a strong association with hypoxia/reoxygenation and sleepiness, as discussed above for memory function, whether hypoxia/reoxygenation can induce irreversible sleepiness must be explored in animal models, without the confounds of obesity, diabetes, and other comorbidities.

E. Effects of Hypoxia/Reoxygenation Exposures on Wake Function

Sleepiness has been assessed in mice 2 wk after exposure to 8 wk of IH and resulted in marked reductions in the average sleep latency across the day and reductions in total wake time for 24 h (694). The magnitude of these effects was similar to changes observed in humans with sleep apnea. Most importantly, these wake impairments in mice were not reversible, even after a 6 mo recovery period in normal oxygen conditions (768). The Veasey laboratory (768) has recently identified the groups of neurons implicated in wakefulness (wake-active neurons) injured by intermittent hypoxia. Forty percent of the noradrenergic neurons in the locus coeruleus and dopaminergic wake-active neurons in the periaqueductal grey were lost in this model, and most remaining wake-active neurons in these nuclei showed impaired wake responses (768). In contrast, orexinergic, histaminergic, serotonergic, and cholinergic wake neurons appeared unperturbed. This differential susceptibility was used to determine the mechanisms by which the wake-active neurons are injured by hypoxia/reoxygenation events. The susceptible catecholaminergic neurons showed increased oxidative injury in response to long-term intermittent hypoxia, relative to resistant neurons. Furthermore, a unique feature of the susceptible wake-active neurons is that they contain NADPH oxidase. Inhibition of this enzyme throughout exposure to hypoxia/reoxygenation or transgenic disruption of the enzyme’s activity largely prevents the hypoxia/reoxygenation wake impairments, supporting the importance of this enzyme in the injury to these neurons (768) (see Fig. 18). Whether a similar catecholaminergic wake neural injury is found in humans with OSA should now be advanced through post mortem neuroanatomical studies of wake active neurons in individuals with and without obstructive sleep apnea.

F. Upper Airway Dilator Motoneurons May Also Be Injured in OSA

Clinical studies have identified neural dysfunction and injury in adults with obstructive sleep apnea. Sensory nerve action potential amplitudes are reduced in individuals with OSA, and treatment of sleep apnea partially reverses this defect, supporting the concept that sleep apnea contributes to this neural dysfunction (146). Electromyographic studies of the palatopharyngeus muscle in individuals with sleep apnea show long polyphasic potentials and reduced amplitude at maximum voluntary effort (660). Consistent with functional impairment, histological studies have identified demyelination of motoneurons in resected palatal tissue in OSA (64, 364, 730). The severity of peripheral nerve dysfunction correlates with oxyhemoglobin desaturations in sleep apnea (397). Whether OSA, independent of obesity, diabetes, and other comorbidities, injures peripheral neurons requires study in animal models. Long-term exposure to hypoxia/reoxygenation impairs hypoglossal whole nerve responsiveness to both glutamatergic and serotonergic excitation of hypoglossal motoneurons (694).

There is increasing evidence that the endoplasmic reticulum (ER) plays a central role in both adaptive responses to and injury from ischemia-reperfusion challenges (49, 126, 228, 229). The unfolded protein response (UPR) in the ER represents an adaptive response to minimize accumulation of misfolded proteins that would be

![Fig. 18. Proposed model of NADPH oxidase injury from hypoxia reoxygenation. Hypoxia/reoxygenation events increase the production of angiotensin II peripherally or in astrocytes, resulting in activation of angiotensin 1A receptors on catecholaminergic neurons. AT receptor activation upregulates NADPH oxidase activity, resulting in oxidative injury. Sleep apnea and intermittent hypoxia are associated with marked inflammation in the brain including inducible nitric oxide synthase (iNOS), cyclooxygenase-2 (COX2), and tumor necrosis factor-α (TNF-α). Whether this proinflammatory response occurs in neurons or adjacent microglial cells should now be advanced.](http://physrev.physiology.org/)

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toxic to the cell. This is accomplished by reducing overall protein translation, increasing the production of chaperones, upregulating clearance of improperly folded proteins, and increasing antioxidant capacity (750). Several components of this protective response are mediated by phosphorylation of eIF-2α. However, when this stress is insurmountable, the ER may take on the role of executioner, activating proapoptotic proteins, including CHOP/GADD153 and caspase-7 (199, 632).

We predicted that IH might activate the UPR and that this response might be injurious in select motoneurons. IH for 8 wk induces significant ER stress in select upper airway motoneurons, including the facial and hypoglossal motoneurons and sparing motor trigeminal (768a). This differential susceptibility was then used to identify the molecular basis of IH susceptibility. Here motoneurons with higher ER stress at baseline develop uncompensated ER stress with exposure to long-term intermittent hypoxia, and many of these motoneurons succumb to apoptosis (768a; Fig. 19).

In summary, intermittent hypoxia, modeling oxygennation patterns of moderate-severe sleep apnea, injures select populations of neurons, including hippocampal, catecholaminergic wake-active, and hypoglossal and facial upper airway motoneurons. It is time to translate these findings to human studies, examining brain tissue from individuals with and without OSA who have died suddenly without a specific neurological diagnosis. As an initial exploration, a focus on the above groups is justified. In light of the mechanisms uncovered in wake and motor neurons, activation of these pathways should also be examined in humans to begin to identify promising therapeutic avenues for the prevention and possibly partial reversal of these injuries.

VIII. FUTURE DIRECTIONS

Substantial advances in several areas of physiology have been made over the past two to three decades, driven by the need to understand the causes, consequences, and treatment of sleep apnea. Major advances in this regard include the neurochemical regulation of upper airway motor neurons and airway caliber; the causes of long-lasting alterations in cardiovascular, biochemical, and neuronal structure and function elicited via intermittent hypoxemia; and the vital importance of the wakeful state on the one hand and variations in sleep state on the other on the regulation of respiratory stability. We close our review of the pathophysiology of sleep apnea by suggesting a few outstanding, fundamental questions for further research.

An experimental approach is needed to provide a more rigorous evaluation of the true prevalence of clinically significant sleep-disordered breathing, than the widely disparate estimates currently provided via epidemiological, correlational approaches. Intervventional treatment trials need to be conducted, preferably at the earliest detectable stages of sleep apnea. Also, prospective trials designed to test the effects of mild to moderate levels of intermittent hypoxia on cardiovascular outcomes and daytime neurocognitive functions need to be...
applied, using the methods currently available in both animals and humans (see sect. v).

How can we best prevent sleep state effects specifically on upper airway collapsibility via pharmacological targets? Do we now have sufficient evidence in support of targets with cholinesterase inhibitors? Do we need to further identify key receptor subtypes?

Obesity complicates defining a causal relationship between OSA and insulin resistance. How can we best design large-scale, controlled studies to determine CPAP treatment effects on insulin resistance and type II diabetes in carefully phenotyped subjects?

Can we identify, on the basis of genotype or phenotype, OSA patients most at risk for development of cardiovascular disease? Can pharmaceutical interventions be used, alone or in combination with traditional therapies, to prevent or readdress the cardiovascular consequences of OSA?

Alternative treatments aimed at minimizing respiratory control system loop gain and breathing instabilities need to be tested specifically in those OSA patients with upper airways that are only moderately susceptible to collapse.

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