Obstructive sleep apnea-hypopnea syndrome: Etiology and diagnosis

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INTRODUCTION

Sleep disordered breathing is a common chronic condition in the general population characterized by repeated episodes of apnea and hypopnea during sleep. It can present as obstructive sleep apnea/hypopnea (OSAH) disorder, central sleep apnea (CSA), or mixed sleep disordered breathing. When OSAH disorder is associated with daytime sleepiness, it is called obstructive sleep apnea-hypopnea syndrome (OSAHS).

Historically, sleep apnea was first described in the early nineteenth century when Charles Dickens reported in his book “The Posthumous Papers of the Pickwick Club” the obese young man that was sleepy and snoring “Said the old gentleman, he’s always asleep. Goes on errands fast asleep, and snores as he waits at table”. Since then the Pickwickian syndrome, known as obesity hypoventilation syndrome, is defined as a combination of obesity, snoring, and excessive sleepiness associated with hypoventilation resulting awake hypercapnia. OSAHS was not recognized as a clinical disorder until nearly 100 years later.

How common is OSAHS?
OSAHS is a common chronic disorder and the most common of all sleep disorders. It can occur on a similar frequency as type-1 diabetes and twice that of asthma. It is estimated from the Wisconsin cohort that the prevalence of OSAHS in the United States of America is 9–24% for men and 4–9% for women who were not obese (body mass index <30 kg/m²) and aged 30–60 years old.[1] Although epidemiological data from the Arab countries are lacking, especially related to OSAHS prevalence, it is estimated that millions of patients suffer from OSAHS in the Middle East and Arab countries [Table 1].

What causes OSAHS?
The pathogenesis of sleep disordered breathing involves an interaction between unfavorable pharyngeal anatomy and ventilatory control instability.[2-5] Obstructive sleep apnea is due to anatomic factors that promote pharyngeal narrowing including large neck circumference, cervical

Table 1: Sleep apnea in the Middle East from US Census Bureau, International Data Base, 2004

| USA | Syria* | KSA | UAE |
|-----|--------|-----|-----|
| Population | 293,655,405 | 18,016,874 | 25,795,938 | 2,534,915 |
| OSA prevalence | 19,433,078 | 1,192,293 | 1,707,084 | 167,023 |
| OSA rate | 1,192,293 | 36,0337 | 51,5918 | 50,478 |

USA - United States of America; KSA - Kingdom of Saudi Arabia; UAE - United Arab Emirates. (*Extrapolated statistics)
soft tissue, vessels, and bony structures. Many of these factors promote pharyngeal collapsibility by decreasing the caliber of the upper airway or by increasing the upper airway surrounding pressure. Increased upper airway collapsibility during sleep has been linked to structural changes in the surrounding boney and soft tissues, which is best measured by determining the critical collapsing pressure or compliance under inhibited neuromuscular activity (passive Pkrit or passive Cua, respectively).

CSH is due to an absent or reduced ventilatory motor output, which is an important determinant of upper airway patency during sleep, especially in individuals with high susceptibility to pharyngeal collapse such as patients with sleep apnea and snoring individuals. When ventilatory motor output oscillates during periodic breathing, pharyngeal narrowing or obstruction occurs at the nadir of ventilatory motor output, especially in individuals with a highly collapsible airway. Recent studies observed that when airflow obstruction occurred during sleep, it triggered an overshoot of ventilation with or without arousals from sleep. The events followed by arousals, however, had greater overshoot ventilation and ensuring obstructive events than those without arousals. These findings indicate that when neuromuscular responses to airflow obstruction fail to compensate, it trigger breathing instability and perpetuate recurrent apnea and hypopneas. Nevertheless, the mechanism of pharyngeal narrowing or obstruction during sleep remains unclear.

Pharyngeal occlusion during obstructive apnea is often described as an inspiratory phenomenon, caused by negative collapsing pressure. However, several lines of evidence indicate that expiratory narrowing may be a significant contributor to pharyngeal obstruction during sleep. More recently, it was found that reduced ventilatory motor output leads to pharyngeal narrowing during expiration. The magnitude of expiratory narrowing correlated with body mass index, indicating that neuromuscular and anatomic factors contribute to upper airway patency during NREM sleep.

To ascertain the mechanisms of pharyngeal narrowing during sleep, it is useful to consider the hypotonic pharynx during apnea or hypopnea as a Starling Resistor with a collapsible segment governed by the principles of flow through collapsible tubes [Figure 1]. Accordingly, if the upstream pressure is below the critical closing pressure (Pcrit), the collapsible segment is closed, and no flow occurs (i.e., apnea). Flow commences when upstream pressure exceeds Pcrit and is determined by the gradient between the upstream segment and Pcrit. The critical closing pressure reflects the surrounding tissue pressure (extraluminal pressure) being negative in normal subjects and positive in patients with OSA. The aforementioned principles operate during both phases of respiration because Pcrit and upstream pressure may change dynamically throughout the respiratory cycle. Upstream pressure is the mask pressure during inspiration and the supraglottic pressure during expiration. Thus, the effects of decreased ventilatory motor output on upper airway patency may differ by the phase of respiration. Decreased ventilatory motor output diminishes the magnitude of negative supraglottic airway pressure, hence increasing downstream pressure with no adverse effects on inspiratory upper airway patency. In contrast, supraglottic pressure is the upstream pressure during expiration and is a major determinant of expiratory flow. Consequently, expiratory pharyngeal obstruction would occur if the supraglottic pressure falls below Pcrit. This may explain the occurrence of pharyngeal occlusion during central apnea or periods of very low drive (as depicted in Figure 2).

What predisposes to OSAHS?

A spectrum of risk factors predispose to OSAHS as depicted in Figure 3. The condition is more common overweight men. In fact obesity is considered the most important risk factor for OSAHS. Obesity as defined by body mass index (BMI) equal or greater than 30 kg/m² has become a global epidemic throughout the world. According to the World Health Organization estimation performed in 2005, more than 30% of the general population in the United States has obesity. This high prevalence is now seen in countries in Europe and the Middle East such as Syria, Saudi Arabia, and...
United Arab Emirates [Table 2]. This increased prevalence of obesity is strongly linked to higher incidence of SDB and OSAHS; therefore screening this type of population for early detection of morbid conditions is vital in the clinical assessment by any clinician or health care provider. Other risk factors include chronic heart failure patients (estimated prevalence of 40–80%),[10] cervical spinal cord injury patients (estimated prevalence of 60%),[16] and stroke (estimated prevalence of 44–72%).[20-23]

Why is it important to recognize OSAHS as early as possible?
Many people who have sleep apnea do not realize it. Drowsiness and lack of concentration contribute to increased traffic accidents.[24] OSAHS can lead to adverse outcomes on the public health, including hypertension, cardiovascular, metabolic, and neuropsychological consequences.[1,25-28] Hypopneas comprise a significant component of OSAHS and may also cause deleterious cardiovascular consequences, particularly when accompanied by desaturation (>4% from baseline).[27] More recently, it has been found in well-designed longitudinal study that moderate and severe sleep apnea are associated with three-fold increase in risk of ischemic stroke after 8 years of follow-up of community-dwelling men and women.[29,30]

OSAHS has been associated with several neuropsychological impairments thought to be due to structural changes in the brain.[31] Recurrent apnea and hypopnea events during sleep result in intermittent hypoxia, hypo-, and hypercapnia and sleep fragmentation. Intermittent hypoxia, hypo-, and hypercapnia are associated with altered vasomotor protection in the central nervous system which can contribute to the structural changes in the brain. A recent study demonstrated that cognitive impairment is associated with decreased gray-matter volume in specific areas of the brain such as frontal and hippocampal regions which associated with decreased executive function and short-term memory. Interestingly, these cognitive and structural changes were reversed after 3 months of OSA treatment.[31] Thus, early recognition of sleep disordered breathing is important step toward effective prevention of such serious consequences in near future.

What are the clinical manifestations of OSAHS?
OSAHS is commonly associated with snoring and excessive daytime sleepiness. Snoring is the most common nocturnal symptoms that could be loud and disturbing to the bed partner’s sleep. Bed partners may report also cessation of breathing (witnessed apnea) which ends by snorting sound or gasping for air. Other nocturnal symptoms are unexplained awakenings, nocturia, and restless sleep. During the day, patient commonly complain of morning headache, not refreshed from sleep, memory problems, and difficulty concentrating.
OSAHS should be suspected clinically by presenting history, physical findings (as delineated in Table 3), and then confirmed by specific sleep tests. The severity of OSAHS can range from mild snoring to hypopnea and sleep apnea. Apnea is usually characterized as cyclical cessation of breathing (for a duration of at least 10 sec), which can be central, obstructive, or mixed in its etiology. Hypopnea is defined as a reduction in airflow resulting short period of awakening (arousal) or decreased oxygenation (destauration of 3–4%).

When to refer patients for evaluation?
Patients who report excessive daytime sleepiness should be asked specific questions to assess the severity of this sleepiness and whether it occurs in situations that require alertness such as during driving or at work. Epworth sleepiness scale is the most common questionnaire used in the clinical settings to assess the level of sleepiness and to follow it objectively. In specific patient phenotypes such as stroke, spinal cord injury, or heart failure, there is dissociation between OSAHS from hypersomnia and obesity. Therefore, in such patients a high index of suspension coupled with meticulous evaluation and clinical history followed by screening study may be necessary as early as possible.

In general, OSAHS should be suspected by the health care provider in the following cases:
• Loud snoring
• Obesity: BMI >40 kg/m²
• Excessive daytime sleepiness
• Stroke patients
• Neuromuscular disorders (such as spinal cord injury or Parkinson’s disease).

How is the diagnosis made?
The diagnosis of sleep disordered breathing is confirmed by an overnight sleep study or polysomnography that is usually done in the sleep laboratory or the patient home. During laboratory-based polysomnography, multiple physiological parameters are measured and recorded simultaneously with the sleep stages measured by a standard electroencephalogram. The respiratory parameters include a flow channel (usually a thermistor placed on the nose and mouth), effort channels (such as respiratory impedance plethysmography), and pulse oximetry. The analysis of the thermistor signal alone cannot accurately score hypopneic events and that nasal pressure analysis measured via nasal prongs connected to a pressure transducer is a simple and useful way to identify nocturnal breathing abnormalities. Other signals that are measured also include electromyogram from the legs and chin, electrocardiogram, and snoring channel. Sleep disordered breathing can be diagnosed also by in-house polysomnography portable monitors (PM). PM are usually divided into three categories: Level II, full polysomnography; level III, at least four channels (flow, effort, oximetry, and ECG); level IV, less than four channels (including oximetry). There is now strong evidence that level III portable monitoring can confirm or exclude the diagnosis of OSAHS with absolute differences between the home and laboratory respiratory disturbance index less than 10 for 65% of patients.\(^{[32]}\)

The 2007 Clinical Guidelines advised that PM should be used in patients with a “high pretest probability” for OSA.\(^{[27]}\) PM is not appropriate for the diagnosis of OSA in patients with significant comorbid medical conditions, for the diagnosis of comorbid sleep disorders, or for general screening of asymptomatic populations. The guidelines also stated that PM may be indicated for the diagnosis of OSA in patients for whom in-laboratory PSG is not possible. It is required that PM must record airflow, respiratory effort, and blood oxygenation. The airflow, effort, and oximetric biosensors conventionally used for in-laboratory PSG should be used in PM. The AASM task force recommendations for PM use are summarized in Table 4. In highly suspected cases, the utility of a diagnostic algorithm in conjunction with autoadjusting positive airway pressure titration in the

| Table 3: Clinical presentation |
|-------------------------------|
| History                      | Exam                                |
| Night time                   | Obesity: BMI and Neck               |
| Snoring                      | HENT:                               |
| Gasping for air              | Craniofacial abnormalities          |
| Witnessed apnea              | Glossomegaly                        |
| Restless sleep               | Enlarged tonsils (I-IV)             |
| Day time                     | Mallampati classification            |
| Excessive sleepiness         | Lateral narrowing                   |
| Unrefreshed on waking        | Retrognathia                        |
| Morning headache             | Nasal congestion or polyps          |

BMI - Body mass index, HENT - Head, neck, nose and throat

1. Performed under the auspices of an AASM-accredited comprehensive sleep medicine program.
2. An experienced sleep technologist must apply the sensors or directly educate patients.
3. The PM device must allow for display of raw data with the capability of manual scoring or editing of automated scoring.
4. A board certified sleep specialist, or an individual who fulfills the eligibility criteria for the sleep medicine certification examination, must review the raw data from PM using scoring criteria consistent with current published AASM standards.
5. A follow-up visit to review test results should be performed for all patients undergoing PM.
6. Negative or technically inadequate PM tests in patients with a high pretest probability of moderate to severe OSA should prompt in-laboratory polysomnography.
initial management of obstructive sleep apnea has been reported in one study to be equivalent to conventional polysomnography.\[33\]

This approach can be used alternatively in areas with limited resources such as the Middle East. It should be noted, however, that when the test is negative with high clinical suspicion, a full polysomnography should be performed to rule out the disease.

**What is the treatment for OSAHS?**

The treatment of OSAHS is beyond the scope of this review. Nevertheless the following management strategies are available: Lifestyle changes, mouthpieces, breathing devices, and surgery are used to treat OSAHS. The goals of treating sleep apnea are to relieve symptoms such as loud snoring and daytime sleepiness and to restore normal breathing during sleep. Lifestyle changes (weight loss and exercise), positional therapy, and mouthpieces can be used to treat mild sleep apnea. People who have moderate or severe sleep apnea will need breathing devices, such as nasal continuous positive airway pressure or surgery.

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