Snoring-Induced Nerve Lesions in the Upper Airway

*Rajesh P Poothrikovil and Mohammed A Al Abri

The prevalence of habitual snoring is extremely high in the general population, and is reported to be roughly 40% in men and 20% in women. The low-frequency vibrations of snoring may cause physical trauma and, more specifically, peripheral nerve injuries, just as jobs which require workers to use vibrating tools over the course of many years result in local nerve lesions in the hands. Histopathological analysis of upper airway (UA) tissues (e.g. tonsils and adenoids) supported by the American Academy of Sleep Medicine (AASM) defines snoring as a sound originating from the upper airway (UA) that does not occur with apnoea or hypventilation, and that is caused by vibrations of different tissues in the pharynx. A person who snores for more than 10–20% of a monitored night, or more than 3 or 4 nights a week is classified as a habitual snorer. An association between snoring and obstructive sleep apnoea syndrome (OSAS) was first observed in 1975. OSAS is considered a progressive disorder that starts, often early in life, with habitual snoring. Based on different epidemiological studies performed between 1980 and 2007, the mean prevalence of snoring in the general population is approximately 22% in men and 37% in women; however, the prevalence of obstructive sleep apnoea (OSA) is 4% in men and 2% in women. OSAS involves the intermittent cessation of breathing due to UA obstruction. In a person with OSAS, the UA muscles relax excessively during sleep. This allows tissues (e.g. tonsils and adenoids) supported by the

Department of Clinical Physiology, Sultan Qaboos University Hospital, Muscat, Oman
*Corresponding Author e-mail: rajeshthrissur@yahoo.com
UA muscles to be drawn into the airway with each inspiration, and thereby obstruct the airflow. The sufferer makes increasingly stronger respiratory efforts to overcome the blockage, but ultimately arouses briefly to take several fast, deep breaths. During the arousal, the sufferer may gasp or snore loudly. This whole cycle recurs on resumption of sleep. If left untreated, OSAS leads to excessive daytime sleepiness, cognitive dysfunction, and impaired work performance, and is detrimental to the sufferer’s health and quality of life. Clinical research shows that OSAS is strongly linked to a range of serious and even life-threatening chronic diseases such as stroke, heart failure, hypertension, diabetes, obesity and coronary heart diseases.

The aetiology of OSAS is only partly known. Sleep disordered breathing (SDB) results from an imbalance between negative pharyngeal pressure and the opposing force of the UA muscles. Abnormal function of the UA, such as sleep-related suppression of UA muscle activity and a decrease in UA dilator muscle force are thought to be the main source of obstructions and symptoms in the OSAS.15,16 Phasic activation of the muscles of the nose, pharynx and larynx has been shown to occur before diaphragm and intercostal muscle activity, suggesting a pre-activation of the UA muscles in preparation for the development of negative pressure.17,18 The UA is rich in neural receptors which is a key factor of tonic genioglossus electromyogram (EMG) activity. During inspiration, the UA dilatory muscles, the genioglossus and geniohyoid contract, or shorten, their muscle fibres through an increase in EMG activity. These inspiration-related muscle activations result in the enlargement of the UA.19 Loss of genioglossus EMG tone may lead to an increase in pharyngeal resistance.20 In order to compensate for abnormal anatomy and/or a more collapsible pharyngeal airway, OSAS patients display augmented genioglossus muscle activity during wakefulness as compared with healthy subjects.21 This reflex compensatory neuromuscular mechanism is lost at sleep onset (loss of awake compensation) in both the control and in OSAS patients, but is associated with pharyngeal collapse under conditions of chronic airway loading.22 When the tongue moves posteriorly due to a decrease in genioglossus muscle activity, the base of the

**Figure 1:** Anatomy of the upper airway showing the main segments: nasopharynx, velopharynx, oropharynx, and hypopharynx.14

**Anatomy and Physiology of the Upper Airway during Sleep**

The UA is a complicated structure, usually divided into four anatomical subsegments: the nasopharynx, between the nares and hard palate; the velopharynx, between the hard palate and soft palate; the oropharynx, from the soft palate to the epiglottis, and the hypopharynx, from the base of the tongue to the larynx [Figure 1]. This total structure forms a passageway for movement of air from the nose to the lungs, and also participates in many competing physiological functions such as phonation and deglutition.9 The UA is surrounded by 20 or more muscles, known collectively as UA muscles, which actively constrict and dilate the UA lumen.10 These muscles can be broadly classified into four groups: muscles regulating the position of the soft palate, tongue, hyoid apparatus, and the posterolateral pharyngeal walls. These muscle groups interact in a complex fashion to determine the patency of the airway. The mandible and hyoid bones are the main craniofacial bony structures that determine the airway’s size.11 As there is no rigid structural support, the shape and size of the UA is dependent upon the position of soft tissue structures, like the soft palate, tongue, and the walls of the oropharynx. Radiographic images show that the narrowest region in both patients with OSAS and non-obese controls, while awake, is the region posterior to the soft palate.12,13 The cross sectional area of this retropalatal region is smaller in OSAS patients than in non-obese controls.
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takes multiple physiological measurements and is an excellent tool for evaluating sleep disorders. Snoring can be detected and evaluated by means of small sensors in the form of piezo crystals or dynamic microphones taped laterally to the thyroid cartilage, or by utilising a nasal pressure transducer which continuously samples nasal air turbulence through a cannula. PSG provides an opportunity to assess the consequences of SDB, such as nocturnal desaturation, frequent arousals, and excessive day time sleepiness, and also plays an important role in the titration of continuous positive airway pressure (CPAP) treatment.

The palatopharyngeal muscle is of anatomical importance in the pharynx as it forms the internal longitudinal muscular layer around the wall of the pharynx. The muscle is located at the major site of obstruction in patients with OSAS and is therefore exposed to the vibration and stretch induced by heavy snoring and obstructive breathing. It has been reported that long term employment which involves the use of vibrating tools such as jack hammers can cause local nerve lesions in the hands. Researchers observed pathological changes in the vibration-exposed fingers, including a marked loss of nerve fibres suggestive of demyelinating neuropathy in the peripheral
nerves. Similarly, snoring produces a low-frequency vibration which may also cause peripheral nerve injury and physical trauma.

Snoring-induced Upper Airway Nerve Damage

THERMAL SENSITIVITY

Larsson et al. reported an impairment of thermal sensitivity in the oropharynx of patients with OSAS as compared with non-snoring age-matched control subjects. Some patients with OSAS were completely unable to differentiate between heat and cold while tested on the tonsillar pillars, whereas no differences was noted at the tip of the tongue in patient and control groups, which indicates a local sensory dysfunction. They postulated that snoring-related vibrations and the deformation of UA structures in the case of apnoeas could lead to a very local pharyngeal sensory neuropathy, which could contribute to UA dysfunction during sleep. As there were only OSAS and non-snoring control groups in this study, a comparison between apnoeic snorers and non-apnoeic snorers was not possible.

VASODILATION

In another study, laser Doppler perfusion monitoring combined with electrical stimulation (a method used to test vascular reactivity) was performed in the mucosa of the soft palate in patients with various degrees of UA obstruction and control subjects. Habitual snorers and patients with mild OSA showed exaggerated vasodilation as compared to controls. This could be the result of minor lesions with consequent re-innervation which increased the sensitivity to mechanical stimuli. In contrast, patients with severe OSAS showed significantly reduced vasodilation as compared to controls, which could be due to the almost complete loss of afferent C-fibres, representing a permanent injury. These disturbances in the micro-circulation indicate the presence of a local afferent nerve lesion with a progressive nature in heavy snorers, both those with and without OSAS.

SENSORY FUNCTIONS

Kimoff et al. assessed sensory functions of the UA (oropharynx) using two-point discrimination and vibration sensory threshold methods in snorers and OSAS patients, and compared with age-matched controls. They also assessed two control points (lower lip and hand) for each subject. This study showed that the sensory detection threshold using both methods was significantly increased in snorers and OSAS patients as compared to controls. Importantly, there was no significant difference between snorers and patients with OSAS. In contrast, the sensory threshold at control sites was similar in patients and controls. Vibrometry is usually employed to assess the functional integrity of the largest afferent sensory fibres to diagnose polyneuropathy. These findings reinforce the presence of a selective impairment of UA mucosal sensory function in patients with OSAS and in those who are heavy snorers. They suggest that this impairment may be an early change in the progression of UA obstruction during sleep, possibly developing as a consequence of vibration-related oedema or neural damage during snoring. Interestingly, after 6 months of continuous CPAP usage, OSAS patients showed a significant improvement in vibration sensory thresholds thus providing strong evidence of the effectiveness of CPAP treatment. These results increase the importance of CPAP usage in the field of sleep medicine as an effective and non-comparable treatment. However, CPAP treatment fails to improve two-point discrimination, indicating a degree of permanent injury. Beginning CPAP treatment before symptoms worsen could be useful in limiting the severity of this permanent neural injury to the pharyngeal muscles.

HISTOPATHOLOGIC FEATURES

Woodson et al. analysed the histopathologic features of pharyngeal tissue. Transverse sections of the distal soft palate and uvula were qualitatively compared between apnoeics, severe snorers, and non-snorers using light and electron microscopy. Light microscopy of both apnoeics and non-apnoeic snorers revealed similar abnormalities such as mucous gland hypertrophy, focal atrophy of muscle fibres, and extensive oedema of the lamina propria with vascular dilation. No distinctive histopathologic findings were associated with the development of apnoea. Electron microscopy was used to reveal frequent focal degeneration of myelinated nerve fibres and axons in severe apnoeics. Similar histopathologic changes were
noted in apnoeics and non-apnoeic snorers. This is an indication of a common aetiology related to snoring-induced vibration trauma to pharyngeal tissue rather than directly related to apnoea or desaturation.

DENERVATION

Friberg et al. described a proliferation (increased density of sensory nerve terminals with abnormal localisation and appearance) of nerve endings in a pattern suggestive of nerve injury in biopsy specimens from the oropharyngeal mucosa of some snorers. However, the study focused mostly on OSAS subjects. No such abnormalities were detected in non-snoring control subjects. Other muscle biopsy studies illustrated similar denervation type changes in palatopharyngeal muscles. The relationship between vibration and stretch-induced trauma was reinforced in a study which showed a significant increase of morphological abnormalities characteristic of neurogenic lesions (type grouping, fascicular atrophy, and grouped atrophy) in the palatopharyngeal muscle of the entire group of snoring patients as compared to non-snoring controls. They observed that the degree of muscle pathology increased in parallel with the proportion of obstructive breathing during sleep. This indicates that the proportion of the total sleep time spent in periodic obstructive breathing may be a surrogate quantitative measure of the magnitude of the snoring trauma to the pharyngeal tissues. Another finding in this study is that the vibration trauma of habitual snoring itself could initiate a local neurogenic lesion in vulnerable patients, before the additional trauma of stretch caused by periodic obstructive breathing. Palatopharyngeal muscle hypertrophy noted in some snorers could be neurogenic and could be due to the chronic stretching of innervated and denervated muscle fibres and/or overuse of partially denervated muscles.

NEUROPHYSIOLOGICAL ASSESSMENTS

Finally, in a recent study, Svanborg conducted neurophysiological assessments of UA muscles by recording concentric needle EMG activity of palatopharyngeal muscles in 12 OSAS patients. Ten out of 12 cases showed reduced EMG activity (recruitment pattern) at maximal voluntary effort with long and polyphasic (increased duration) motor unit potentials. These features typify motor neuropathy (chronic denervation and reinnervation). Two patients showed spontaneous denervation activity (fibrillations and positive waves) suggestive of an active neuropathic process. Such findings were present in only 3 out of 15 habitual snorers. This study was of great importance in that it gives strong electrophysiological evidence of motor neuropathy in the UA muscles of OSA patients. These findings reinforce the causative role of peripheral neurogenic lesions, in the progression from habitual snoring to clinical OSAS.

Upper Airway Neuropathy in Progressive Snorers’ Disease

It is obvious that the local UA afferent (sensory) and efferent (motor) nerve lesions are present due to snoring-induced vibration trauma in some patients with snoring and in most patients with OSAS. It has been reported that the higher sound intensity in apnoeic snorers versus non-apnoeic snorers is a result of greater negative pressures on the resumption of breathing, resulting in high flow rates, turbulent flow, and greater forces on the vibrating structures. This reflex mechanism, as a reaction to the negative intrapharyngeal pressure, is responsible for the activation of dilatory muscles. Numerous studies have described protective UA dilator reflex responses to pulses of negative airway pressure, which act to maintain airway patency during sleep. This reflex is assumed to be mediated by intra- or submucosal mechanosensory receptors. The neural drive to the UA dilator muscles is integrated at the level of brainstem motor nuclei where multiple neural inputs combine to produce a unitary output to the muscles. Neurogenic lesions affecting upper airway muscles impair the ability of its reflexogenic dilation, leading to increased possibility of UA collapse. A reduced pharyngeal lumen and other risk factors (i.e. obesity and anatomic abnormalities) lead to an increased degree of periodic obstructed breathing. This causes more morphological abnormalities and neurogenic lesions, which supports the hypothesis of the progressive nature of snorers’ disease.
Causes and Treatment Options for Snoring

Other causes of habitual snoring, such as nasal septal deviation, nasal valve obstruction, chronic nocturnal nasal congestion, and craniofacial abnormalities should be well identified. Patients with chronic nasal obstruction often struggle to tolerate nasal CPAP. Humidification of inhaled air, correction of potential leakage of the nasal mask, and a trial of an oro-nasal mask are initial steps that can be considered to acclimatise patients to the CPAP treatment. Patients who still cannot tolerate CPAP, or have obvious nasal polyps or a distinctive abnormality of the nasal anatomy, should consider surgical treatment. Correction of nasal obstructions has been reported to be an effective treatment of OSAS in patients who have nasal obstructions, but in those with craniofacial abnormalities, correction has proved ineffective. Improving nasal patency by external nasal dilators has some beneficial effects on subjective snoring, but not in patients with apnoeas. Current evidence suggests that the nose may not play a significant role in the pathogenesis of OSA, but it seems to be of some relevance in the origin of snoring.

Obesity is an independent risk factor for the development of snoring. Simple snoring can be treated with general measures, including weight control and loss, avoidance of detrimental habits and toxic substances that interfere with sleep, modification of sleeping position and physical exercise. Advancement of the mandible using a dental appliance has been shown to reduce the severity of OSA. It is an effective alternate to CPAP in the treatment of snoring and milder form of OSAS. Variable mandibular advancement devices have also been successful in the treatment of patients with severe OSAS. Because the tongue attaches to the mandible directly, the tongue is expected to move anteriorly with the forward displacement of the mandible, thus improving oropharyngeal airway patency. However, no studies have been published to prove the efficacy of mandibular advancement devices in the improvement of upper airway neural injury in snoring patients.

Conclusion

In summary, the progressive nature of OSAS can be partly explained neurologically. OSAS begins with snoring-induced vibration trauma of nerves associated with the UA muscles. Histopathological, neurological, and neurophysiological signs of nerve lesions are detected in simple snorers without obstructive episodes. Periodic obstructive breathing phases expressed as a fraction of total sleep time could be a quantitative measure of the magnitude of snoring trauma to the pharyngeal tissues. Subjects with habitual snoring at risk of development of OSAS should be identified early. Beginning CPAP therapy before symptoms worsen should be considered as a preventive measure against permanent UA neuronal injury. Other options such as weight loss in obese individuals, surgical intervention for nasal obstructions, and mandibular advancement devices for craniofacial anomalies and narrow airway can also be considered in suitable patients, especially those who cannot tolerate CPAP. Studies of novel techniques may help in developing protocols to assess and manage habitual snorers.

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