Obesity is the most important risk factor for OSA, although people who aren't overweight can have OSA. Obesity increases the risk for sleep apnea because fatty tissue in breathing passage reduces the space for air to pass through.\(^1\) There is a dose-dependent relationship, with higher BMI indicating higher risk. A moderate to severe OSA category, defined by an Apnea-hypopnea Index (AHI) \(\geq 15\), was independently associated with increased BMI, neck circumference and waist circumference.\(^1\) Recently, patients with Diabetes Mellitus Type 2 and higher BMI were shown to be more likely to have severe OSA.\(^4,15\)
Case Report
A 65-year-old male with a long-standing history of moderate snoring noted that, in recent years, the snoring had worsened so much that his wife banned him from their bedroom. Since his retirement at 60, due to reduced physical activity he gained 20 kg and started suffering from knee problems. He noted increased fatigue, daytime sleepiness and some trouble in concentrating. His current BMI=51.2 Kg/sq. m. Arterial blood pressure was 120/75 mmHg, pulse was 95 bpm, respiratory rate was 25/min, and he had tachypnea. On respiratory system examination, there were both inspiratory & expiratory rhonchi in all the lung fields. Other system examinations were normal.

Laboratory examinations were normal except for a mild anemia (Hemoglobin = 11.2 gr) and serum IgE =181 IU/ml. Chest X-Ray and spirometry revealed mild obstructive defect with positive reversibility to exclude an upper airway obstruction. Computed tomography (CT) of the thorax and paranasal sinuses and magnetic resonance imaging (MRI) of the neck, which were obtained to evaluate this, were normal. On ear-nose-throat and laryngoscopy examinations, the upper airway was normal. The patient was evaluated by the gastroenterology department to exclude gastroesophageal reflux disorder. Bronchoscopy tests and transthoracic echocardiography of the patient revealed normal results. He noticed that he had been using meter dose inhaler properly for last 10 years as he was diagnosed as bronchial asthma. He reported that 10 years back he performed a polysomnography as suggested by his pulmonologist and it was normal. He is on medication regimen as treatment for Bronchial Asthma but he otherwise denied having any medical problems. He had a tonsillectomy in his childhood and had no history of thyroid disease.

Physical examination showed nasal congestion with moderately swollen, pale turbinates and no purulent discharge. The septum was midline. Oropharyngeal examination showed no tonsils and a low soft palate with elongated uvula that tended to collapse against the posterior aspect of the pharynx and abutted the base of tongue. Fiber optic laryngeal examination showed a normal larynx with moderate collapse of the lateral pharyngeal walls in “blocked” inspiration (a reverse Müller’s maneuver whereby the patient holds his nose, closes his mouth, and attempts to breathe inward).

He had a near normal neck and was overweight. The working diagnosis was obstructive sleep apnea after scoring with Epworth’s score. A overnight sleep study was performed in our Sleep Lab including Oximetry and he was diagnosed as a case of severe obstructive sleep Apnea with AHI 51.

On next day of sleep test, patient started to use CPAP as per advice. On first night he was very much uncomfortable with the face mask and could use only 30 minutes and next morning come with concern. After proper counselling he went back home and came for follow up after one week. He was happy since he was able to enjoy a good sleep after many years.
General and behavioral measures, such as weight loss, avoidance of alcohol for 4-6 hours prior to bedtime, and sleeping on one’s side rather than on the stomach was advised. The best data suggest that a 10% reduction in weight leads to a 26% reduction in the respiratory disturbance index (RDI).13 Patient was also referred to dietician for proper dietary advice & surgeon for consultation for bariatric surgery.

Discussion
Several risk factors, including obesity, male sex, age and heritable factors, have been associated with an increased prevalence of obstructive sleep apnea in the general population. Among these, obesity is one of the strongest sleep apnea risk factors. Young and colleagues showed that a 1-SD increase in BMI was associated with a four-fold increased risk for prevalent sleep apnea and they have demonstrated a sleep apnea prevalence of approximately 40% in moderately overweight men from the community who are otherwise healthy. In severe obesity (BMI > 40 kg/m²), the prevalence of sleep apnea was estimated to vary between 40% and 90%. In addition, Peppard and colleagues have provided further evidence for a link between sleep apnea and obesity by demonstrating that a 10% change in body weight was associated with a parallel change of approximately 30% in the apnea-hypopnea index (AHI), the major index of sleep apnea severity.

Weight loss remains a highly effective strategy for treating sleep apnea. In two controlled studies, investigators have demonstrated that a 10% to 15% reduction in body weight leads to an approximately 50% reduction in sleep apnea severity (AHI) in moderately obese male patients. In recent years, bariatric surgical procedures have been increasingly used for the treatment of severe obesity.

Obesity is associated with anatomic alterations that predispose to upper airway obstruction during sleep. These alterations may result from adiposity around the pharynx and events as follows. First, increase in neck circumference and fat deposit around the upper airway. Second, upper airway collapsibility is higher in obese compared with non-obese individuals and does not decrease appropriately when the pharynx is dilated by advancing the mandible anteriorly. Third, obesity and especially central obesity have been associated with reductions in lung volume, which leads to a loss of caudal traction on the upper airway and an increase in pharyngeal collapsibility, increasing continuous positive airway pressure requirements and a greater severity of sleep apnea. Thus, obesity imposes mechanical loads on both the upper airway and respiratory system that predispose to upper airway narrowing, collapse and airflow obstruction during sleep. Although central adiposity is associated with structural defects that compromise airway patency, the mechanisms causing these elevations in upper airway mechanical loads in obesity are not well understood.

Conclusion
Improvements in sleep apnea with weight loss have been related to effects of adiposity on upper airway function during sleep. In controlled weight loss intervention studies, we demonstrated decreases in upper airway collapsibility during sleep with weight loss. Central obesity accounts for the strong male predominance of this disorder, whereas peripheral adiposity may protect women from developing
adiposity may protect women from developing sleep apnea. Obesity and particularly central adiposity can increase sleep apnea susceptibility by increasing upper airway mechanical loads and/or decreasing compensatory neuromuscular responses. These effects may be mediated by circulating adipokines, which influence body fat distribution and CNS activity. As patients with sleep apnea lose weight, improvements in upper airway function and disease severity are likely related to the amount and patterns of weight loss as well as relative changes in protective and pathogenic adipokines.

References