Alteration in respiratory physiology in obesity for anesthesia-critical care physician

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ABSTRACT

Obesity is known to be a major risk factor of a whole range of cardiovascular, metabolic and respiratory disorders. The pattern of regional fat distribution plays an important role in the predisposition of obese subjects to respiratory complications. Obesity is responsible for important changes in respiratory function both during spontaneous breathing as well as during general anesthesia and mechanical ventilation. The most characteristic abnormalities consist of decreased functional residual capacity, reduced expiratory reserve volume, decreased compliance and increased resistance of the respiratory system. Breathing at low lung volume promotes airway closure in the dependent lung zones with consequent gas exchange abnormalities even though lung carbon monoxide-diffusing capacity is normal or increased. Weight loss can reduce many of the alterations in pulmonary function related to obesity.

Keywords: lung volumes, compliance, oxygen consumption, obesity, airway resistance, diffusion capacity, ventilation, perfusion, oxygenation, pulmonary physiology

INTRODUCTION

For several decades, the global prevalence of obesity has been rising dramatically (1, 2). The greatest increase has been noted in the United States. Compared with some European countries, the prevalence of obesity in the United States is three times higher than in France, and one and a half times higher than in the United Kingdom (3). Between 1980 and 2004, the prevalence of obesity in the US more than doubled in adults and more than tripled in children. The greatest relative increase has been in the proportion of individuals with a body mass index (BMI) greater than 50 kg/m². This review describes the mechanisms whereby obesity brings about the functional abnormalities on resting and exercise related respiratory physiology.

Lung mechanics

Obesity decreases total respiratory compliance by as much as two-thirds of the normal value measured in non-obese individuals (4). The decrease in compliance was thought to result primarily from a reduced chest wall compliance associated with the deposition of fat in and around the ribs, the diaphragm and the abdomen. Subsequent investigations in healthy obese subjects revealed higher total respiratory system and chest wall elastance during voluntary
muscle relaxation than during paralysis (5), suggesting that incomplete relaxation may have contributed to lower chest wall compliance reported in earlier studies. Actually, the chest wall compliance is usually normal in obese subjects and the decrease in total respiratory compliance is that of the lung. The reduction in lung compliance in obese individuals is exponentially related to BMI (6). This decrement is the result of increased pulmonary blood volume, closure of dependent airways (10), and increased alveolar surface tension due to the reduction in functional residual capacity (FRC) (7-9).

**Lung volumes and spirometry**

The most common and consistent characteristic of obesity on lung function is a reduction in FRC (Figure 1). This derangement reflects the mass load of adipose tissue around the rib cage and abdomen (11). In contrast, residual volume (RV) is usually well preserved, and the RV-to-total lung capacity (TLC) ratio remains normal or slightly increased (12). As a result, ERV decreases exponentially with increasing BMI, even in mild obesity or overweight due to displacement of the diaphragm into the thorax and increased chest wall mass. ERV reduction is greatest in the supine position. The reduction is often so marked that FRC approaches RV. At that point, regional thoracic gas trapping may take place causing an elevated RV/TLC ratio (13). Total lung capacity (TLC) and vital capacity (VC) decrease linearly with a rising body mass index, however, the changes are small, and TLC is usually maintained above the lower limit of normal. A marked abnormality of lung volumes in mild to moderate obesity should raise suspicion of an underlying intrinsic lung disease or neuromuscular pathology except in those with morbid obesity or those with excessive central adiposity (waist-to-hip ratio ≥ 0.95) (14).

Spirometry is normal in mild obesity. As BMI increases, there is a reduction in expiratory flow and a decrease in FEV1 and FVC (15). The ratio of FEV1 to FVC is preserved and even increased, which is attributed to peripheral airway closure and gas trapping, hence reducing the VC. However, the reduction in FEV1 and FVC is strongly correlated with abdominal obesity. FVC, FEV1, and TLC were found to be significantly lower in subjects with upper body fat distribution or central obesity (16).

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**Figure 1 - Impact of obesity on lung volumes.**

IC = Inspiratory capacity, IRV = Inspiratory reserve volume, TV = Tidal volume, ERV = Expiratory reserve volume, RV = Residual volume, FRC = Functional residual capacity, VC = Vital capacity.
Abdominal obesity is responsible for a reduced FEV1/FVC ratio suggesting an effect of obesity on large airway caliber as well. In addition to these spirometric derangements, tidal volume is reduced in severe obesity, and breathing follows a rapid, shallow pattern (17). This functional change is typically due to the elastic load which can be replicated in normal weight subjects with elastic strapping of the chest (18). As FRC becomes less than the closing volume, airway closure occurs during tidal breathing. Together with alveolar collapse, this leads to decreased ventilation of the lung bases, ventilation-perfusion mismatch, and hypoxemia. For these reasons, both the PaO2 and the alveolar-arterial gradient are related to FRC. The improvement of lung function with weight loss supports the causative effects of obesity on respiratory physiology. Following bariatric surgery, restrictive pulmonary mechanics improves significantly with corresponding increase in FEV1, FVC, and in FEV25–75\%. Additionally, the obstructive lung pattern (FEV1/FVC ratio less than 0.8) tends to normalize (19). In one program, weight loss was accompanied by an improvement of 73 ml in FEV1 and 92 ml in FVC for every 10\% relative loss of pretreatment weight (20).

The effect of obesity on lung volumes and chest compliance can be worsened by anesthesia and muscular paralysis, which is manifested by decreased lung volumes, and higher lung and respiratory system elastance. This deterioration in lung function is more pronounced with abdominal surgeries (21), but it also seen in other types of non-abdominal procedures. Although the post-operative spirometry of obese patients shows a decrease in FEV1, FVC and PEF, the reduction in vital capacity is the most prominent in both abdominal and non-abdominal surgery, and is greater with increasing BMI.

Various methods have been tried to compensate for the effects of obesity on lung function, and they range from positional changes, to adopting altered intra-operative ventilation strategies, to the use of prophylactic BiPAP post operatively (22). The reverse Trendelenburg (RT) position is one measure that seems to improve the respiratory compliance and gas exchange in morbidly obese patients during bariatric surgery.

However, it is not clear yet if the beneficial effect of this position can be replicated to all abdominal and non-abdominal surgeries in obese patients. Similarly, in intubated patients with large abdomen (obesity, distended abdomen or large ascites), a 45° RT position is associated with larger tidal volume and lower respiratory rate compared with 90° position. These results suggest that the reverse Trendelenburg could be the optimal position to be used in obese patients, particularly in intubated ICU patients, or those undergoing or recovering from anesthesia and surgery (21-23).

In addition to body positioning, different ventilation strategies are used to improve respiratory function in obese patients. Increasing PEEP to 10 cm H2O in anesthetized or paralyzed patients significantly reduces elastance of the respiratory system, lung, and chest wall, and improves oxygenation.

In one study that used computed tomography to assess atelectasis in morbidly obese anesthetized patients undergoing gastoplasty, recruitment maneuver with 55 cm H2O inspiratory pressure for 10 seconds, followed by 10 cm of PEEP reduced atelectasis and improved oxygenation, while recruiting maneuver alone without PEEP yielded only a transient reduction of atelectasis that was not sustained 20 minutes later. In addition, 10 cm of PEEP alone did not affect atelectasis. Repeating the recruitment maneuver every 10 minutes, in
addition to PEEP of 10 cm H₂O, had better results in terms of improving respiratory compliance and oxygenation. Whether such ventilation strategies can be applied to medical obese patients in the intensive care unit is to be determined (24-26). The use of BiPAP as a prophylactic measure to improve pulmonary function after surgery has been studied in obese patients after undergoing gastroplasty. The prophylactic use of BiPAP System 12/4 (but not 8/4), during the first 24 hours postoperatively reduces pulmonary dysfunction after gastroplasty, and accelerates reestablishment of preoperative pulmonary function, which is reflected in improved FVC and FEV₁, as well as SpO₂. The BiPAP acts through enhancing the alveolar recruitment during inspiration, while preventing the expiratory alveolar collapse, and thus reducing the postoperative restrictive syndrome (27).

Respiratory muscles/work of breathing
Studies on the respiratory muscles of obese individuals are scarce. Overall, obese subjects demonstrate inefficiency of respiratory muscles, most notably the diaphragm. The maximum inspiratory and expiratory pressures at all lung volumes are lower in obese patients compared to controls, without reaching statistical significance however, except in patients with obesity hypoventilation syndrome (OHS).

The maximal voluntary ventilation, a measurement of respiratory muscle endurance, is reduced by 20% in healthy obese individuals and by 45% in patients with OHS (28). It is suggested that the additional load causes a length-tension disadvantage for the diaphragm due to fiber overstretching placing the diaphragmatic fibers at suboptimal length. Furthermore, analysis of the diaphragmatic electromyogram revealed a persistence of activity into early expiration, the length of which also depended on the degree of obesity. These findings indicate that the diaphragm’s volume-generating function in the obese is reduced, and furthermore the persistence of its activity in expiration serves to attenuate the rate of expiratory flow (28-30).

On a cellular level, obesity with high intake-associated lipid accumulation in muscle interferes with cellular mitochondrial function through the generation of reactive oxygen species (31). These compounds lead to lipid membrane peroxidative injury and disruption of mitochondrial-dependent enzymes resulting in decreased oxidative metabolism. A reduced ability to oxidize fatty acids has also been reported in skeletal muscle of obese individuals both before and after weight loss, which would support an intrinsic abnormality of fatty acid oxidation (32).

After weight loss, there is a significant increase and return to normal reference values, with regard to both the strength and endurance of respiratory muscles, with the latter showing greater increases. This improvement in respiratory muscle endurance is related to increased chest wall compliance and pulmonary volumes, as a consequence of weight reduction (33).

Airway resistance
Obese subjects have an increased total respiratory resistance due to a predominantly increased airway resistance rather than chest wall resistance. However, when airway resistance is adjusted for the lung volume at which the measurements are made, specific airway resistance is within the normal range indicating that the apparent reduction in airway caliber in the obese is attributable to the reduction in lung volumes rather than to airway obstruction (34). However, recent investigations have suggested that the increase in resistance may not be entirely due to reduction of FRC since
differences between obese and non-obese may persist after lung volume adjustment \(35, 36\). The mechanism by which obesity could cause increased airway resistance is not well understood. The possible hypotheses include increased atopic reaction related to an enhanced inflammatory state secondary to obesity \(38\). Some in vitro studies, as well as human studies suggest that lower lung volumes secondary to obesity, lead to a reduction in peripheral airway diameter, which over time causes smooth muscle dysfunction, and causes both airways obstruction and hyper-responsiveness. In addition, leptin has been suggested to be involved in the airway dysfunction associated with obesity, through its pro-inflammatory properties and/or via a direct effect on airways smooth muscles \(37\). However, the data addressing this question have been inconclusive so far and further studies are needed to understand the mechanism of increased airway resistance and responsiveness in obesity \(38-45\).

The effect of obesity on airway hyper-responsiveness (AHR) has been inconsistently demonstrated. Investigators simulated obesity-related lung volume reductions in non-asthmatic subjects by externally mass loading the chest wall and abdomen and documented an augmentation of airway responsiveness to metacholine relative to that of control \(46\). The relation between BMI and AHR has also been reported in The European Community Respiratory Health Survey \(47\). However, the association between asthma and obesity in adults and children so far has failed to show a consistent increase in AHR \(38\). In addition, weight loss programs did not result in substantial change in AHR despite documented improvements in lung function. Therefore, there is a plausible mechanism to explain how obesity is implicated in AHR but is not consistently reproducible in clinical studies.

**Control of breathing**

Although some studies investigating ventilatory drive in simple obesity have demonstrated that the ventilatory responses to inhalation of carbon dioxide (DVE/DPCO\(_2\)) are normal, others have indicated a reduced response particularly in patients with obesity hypoventilation syndrome (OHS)\(48-49\). These abnormalities were initially attributed to the mechanical limitations and decreased chest compliance preventing adequate ventilation. However the anticipated response to CO\(_2\) did not improve in OHS patients following weight loss. Further, Vd/Vt did not correlate with subjects’ resting PCO\(_2\)\(50\). One theory proclaims that the diminished responsiveness may represent an adaptive process sparing O\(_2\) for non-ventilatory demands. Yet there is an inherent problem with using ventilatory responses as a marker of respiratory drive because minute ventilation response to a stimulus may also be influenced by respiratory muscle function and respiratory system mechanics. The mouth occlusion pressure (P0.1) believed to reflect neurogenic drive is twice the normal value in mild obesity and increases normally with CO\(_2\) inhalation. In contrast, the P0.1 response to CO\(_2\) in patients with OHS is half that of subjects with simple obesity \(49\). The fact that OHS subjects can normalize their PaCO\(_2\) by hyperventilation provides supportive evidence that ventilatory control is abnormal in OHS \(51\). Hence, the cumulative data indicates that subjects with simple obesity have an enhanced respiratory drive while the respiratory drive of subjects with OHS is either depressed or inappropriately suppressed.

**Oxygen cost of breathing**

In non-obese individuals, the percentage of cardiac output and total body oxygen consumption (VO\(_2\)) dedicated to respiratory muscle work during quiet breathing is very
small (less than 3%). In contrast, the oxygen cost of breathing is 4 to 10 fold higher than normal among subjects with eucapnic obesity (Figure 2). In one study of obese patients undergoing bariatric surgery, a 16% reduction in mean VO$_2$ in obese patients was observed compared to less than 1% reduction in the non-obese during the transition from spontaneous breathing to positive pressure ventilation. This suggests that morbidly obese patients dedicate a disproportionate high percentage of total VO$_2$ for respiratory work.

Of interest, the obese patients demonstrate a significantly lower VO$_2$ when standardized by BMI that has been attributed to the lower blood flow and metabolic rate of adipose tissue compared with lean body tissue. Nevertheless, the lower VO$_2$ standardized to body size does not ameliorate the detrimental impact of morbid obesity on oxygen consumption. This respiratory inefficiency results in a limited ventilatory reserve that predisposes these patients to respiratory failure in the setting of acute pulmonary or systemic illnesses (52,53).

**Ventilation/Perfusion (V/Q)**
Ventilation in non-obese patients is greatest in dependent lung zones and decreases toward the upper zones; however, this distribution may be reversed in obesity. When lungs ventilation and perfusion were examined in obese subjects with ERV at 21% of predicted, the normal tidal breath predominantly distributed to the upper zones, while perfusion was predominant in the lower lung zones.

In contrast, subjects who had an average ERV of 49% of predicted value had normal ventilation distribution (54). Thus, impairment of the V/Q relationships depends on the location of the excess body weight. Individuals with central obesity seem the most affected. Similar results were reproduced in lateral decubitus position (55). This ventilation/perfusion mismatch results from airway closure in the lungs’ dependent areas of obese patients.

**Diffusing capacity and gas exchange**
The diffusing capacity of obese subjects is usually preserved although studies have reported increased and decreased values.

![Figure 2 - Interplay of respiratory mechanics on oxygen consumption in obese patients.](image_url)
An increased DLCO in obese patients is probably related to increased pulmonary blood volume and flow while a decreased DLCO may result from structural changes in the interstitium from lipid deposition or decreased alveolar surface area. In either scenario, weight loss appears to have little effect on diffusing capacity as DLCO values remained unchanged following surgical or medical treatment (57, 58). Morbid obesity is associated with low arterial pressure of oxygen (PaO₂) and increased alveolar-to-arterial oxygen partial pressure difference (58). These changes are usually more prominent in men than in women secondary to gender differences in waist-to-hip ratio. Arterial pressure of carbon dioxide (PaCO₂) is usually normal in obese patients who do not have obesity hypoventilation syndrome. While the gas exchanges improve with peak exercise, obese subjects have a poor compensatory hyperventilation, resulting in low exercise tolerance and premature termination of exercise (59).

The mechanism by which obesity impairs blood gas exchange and oxygenation is related to lower lung volumes and basilar atelectasis secondary to airway closure and alveolar collapse. Increased airway resistance has little role in impaired gas exchange at rest (normal PaCO₂); it may however play a role in the poor exercise tolerance through increased expiratory airflow limitation and dynamic hyperinflation.

Therapeutic measures that are used to improve lung volumes and decrease atelectasis are also associated with improvement of oxygenation and blood gas exchange. In fact, both PaO₂ and alveolar-arterial oxygen difference were ameliorated when PEEP of 10 cm H₂O was applied to paralyzed and anesthetized postoperative obese patients after abdominal surgery, compared to PEEP of zero. Similarly, the alveolar-arterial oxygen difference was significantly reduced when morbidly obese patients undergoing bariatric surgery were placed in reverse Trendelenburg position. In addition, the application of BiPAP 12/4 improved oxygen saturation when used prophylactically in obese patients for 24 hours after undergoing gastroplasty (56, 60).

**Altered exercise respiratory physiology**

At rest, the baseline VO₂ is approximately 25% greater than the VO₂ for non-obese individuals. Because adipose tissue has a lower metabolic rate than other tissues, peak VO₂ uptake adjusted for true body weight is reduced, however peak VO₂ is usually normal or increased when adjusted for ideal body weight (61). Interestingly, the slope of the VO₂ work-rate relationship is unchanged but it is shifted upward by approximately 6 ml/min/kg of extra body weight for a cycle ergometer. This means that an appropriate peak VO₂ standard reference for an obese subject can be predicted by increasing the standard peak VO₂ from the reference body weight by 6 ml/min for each kilogram greater than the reference weight. Other responses may vary depending on the exercise protocol and severity of obesity. Parameters including peak O₂ pulse (VO₂/heart rate (HR)), and anaerobic threshold are usually normal in mild to moderate obesity (62).

The resting HR is usually elevated, reflecting an increase in cardiac output at rest. With exercise, there is a normal HR-VO2 relationship reflected by a normal HR-VO₂ slope and attainment of the predicted HR with no HR reserve.

It is unusual for obese individuals to demonstrate ventilatory limitation despite the abnormalities imposed on the respiratory system at rest. Because ventilation perfusion relationship normalizes during exercise, dead space ventilation usually responds normally with a decrease toward the normal range with exercise (63).
**Pulmonary vasculature**

Pulmonary artery systolic pressure (PASP) correlates echocardiographically with BMI independently of age, gender, or comorbid diseases. Using echocardiography, PASP \( \geq 30 \) mm Hg and \( \geq 35 \) mm Hg occurred in up to 66% and 36% of obese subjects, respectively. For each unit increase in BMI, the PASP increases by 0.1 to 0.4 mm Hg. The exact mechanism of increased PASP in obesity is not clear, likely related however to increased blood volume. Obstructive sleep apnea and pulmonary capillary spasm secondary to nocturnal hypoxia are other possible contributing factors. However in the absence of right heart catheterization, echocardiographic findings should be interpreted with caution (64).

**CONCLUSIONS**

Obesity affects respiratory physiology in many ways, with significant clinical implications. There are few measures that are shown to improve respiratory function in obese patients undergoing medical or surgical treatment (Table 1).

They include reverse Trendelenburg position, higher PEEP with recruitment maneuvers or pressure control mode in ventilated patients, and prophylactic use of BiPAP after surgery/sedation. Increasing positive end expiratory pressure (PEEP) to 10 cm H\(_2\)O significantly reduced elastance of the respiratory system, lung, and chest wall in obese patients.

Additionally, application bi-level positive airway pressure of (BiPAP) 12/4 improves oxygen saturation in obese patients for 24 hours after surgery.

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117

Respiratory physiology in obesity

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