Dyspnea and Activity Limitation in COPD: Mechanical Factors

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MEASUREMENT OF EXERCISE IN COPD

Dyspnea and Activity Limitation in COPD: Mechanical Factors

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ABSTRACT

Dyspnea and activity limitation are the primary symptoms of chronic obstructive pulmonary disease and progress relentlessly as the disease advances. In COPD, dyspnea is multifactorial but abnormal dynamic ventilatory mechanics are believed to be important. Dynamic lung hyperinflation occurs during exercise in the majority of flow-limited patients with chronic obstructive pulmonary disease and may have serious sensory and mechanical consequences. This proposition is supported by several studies, which have shown a close correlation between indices of dynamic lung hyperinflation and measures of both exertional dyspnea and exercise performance. The strength of this association has been further confirmed by studies that have therapeutically manipulated this dependent variable. Relief of exertional dyspnea and improved exercise endurance following bronchodilator therapy correlate well with reduced lung hyperinflation. The mechanisms by which dynamic lung hyperinflation give rise to exertional dyspnea and exercise intolerance are complex. However, recent mechanistic studies suggest that dynamic lung hyperinflation-induced volume restriction and consequent neuromechanical uncoupling of the respiratory system are key mechanisms. This review examines, in some detail, the derangements of ventilatory mechanics that are peculiar to chronic obstructive pulmonary disease and attempts to provide a mechanistic rationale for the attendant respiratory discomfort and activity limitation.

INTRODUCTION

Dyspnea (i.e., perceived respiratory discomfort) is the primary symptom limiting exercise in the majority of patients with more advanced chronic obstructive pulmonary disease (COPD) and can lead to avoidance of activity with consequent skeletal muscle deconditioning (1–6). In a study of 105 clinically stable patients (FEV₁ = 37% predicted), severe breathing discomfort was the primary symptom limiting incremental cycle exercise in 61%; combined dyspnea and leg discomfort limited exercise in 19%, and 18% stopped primarily because of leg discomfort (3). Patients who stopped exercise primarily because of dyspnea, had greater levels of dynamic lung hyperinflation, greater ventilatory constraints and poorer exercise performance than the minority who stopped mainly because of leg discomfort (3). This frequency distribution of exercise limiting symptoms was almost similar to that found in a study in 125 symptomatic patients with COPD entering a pulmonary rehabilitation program (7) and in a combined group of 353 COPD patients undertaking a symptom-limited incremental cycle exercise (5, 8) (Figure 1). Similarly, in a group of 403 patients with moderate-to-severe COPD who undertook constant work-rate cycle exercise, dyspnea again was the main exercise-limiting symptom in the majority (63%), whereas leg discomfort alone and in combination with dyspnea accounted for 9% and 27%, respectively (5, 8) (Figure 1).

In COPD, exercise limitation is clearly multifactorial and ultimately reflects integrated abnormalities of the ventilatory, cardiovascular, peripheral muscle, metabolic and neurosensory systems in highly variable combinations. However, ventilatory limitation is the dominant contributor to exercise curtailment in more advanced disease and is the main focus of this review. In particular, we will review the evidence that lung hyperinflation provides a mechanistic link between expiratory flow-limitation, dyspnea and exercise intolerance.

Pathophysiology of lung hyperinflation in COPD

The volume of air remaining in the lung at the end of spontaneous expiration (i.e., end-expiratory lung volume (EELV))

Ventilatory responses to activity in COPD

Ventilatory responses to exercise in COPD are increased relative to health. Stimuli for excessive ventilation in COPD include: increased chemo-stimulation as a result of the effects of extensive ventilation-perfusion abnormalities, early metabolic acidosis, feedback from activated mechanoreceptors/metaboreceptors in the working peripheral muscles or any combination of the above listed factors. During physical activity, healthy younger subjects tend to decrease EELV as a result of recruitment of abdominal and expiratory rib cage muscles (10, 11). Active reduction of EELV in this manner assists the inspiratory muscles and ensures that tidal volume expansion occurs within the linear compliant portion of the respiratory system’s pressure-volume relation (Figure 2).

With advancing age, changes in the lungs’ connective tissue matrix can lead to expiratory flow-limitation at higher levels of ventilation and therefore the ability to reduce EELV is curtailed during exercise. Although patients with COPD progressively recruit their expiratory muscles during exercise, EELV usually increases as a consequence of expiratory flow-limitation (3, 12–17). This temporary and variable increase in EELV above its baseline value is termed “dynamic” lung hyperinflation (DH). Thus, under any condition of increased ventilation in flow-limited patients (e.g., exercise or voluntary hyperventilation), inspiratory tidal volume increases and expiratory time diminishes further as breathing frequency increases above the baseline value, causing further acute-on-chronic DH (3, 12, 18–20).

It has been established for some time that DH occurs in flow-limited patients under conditions of increased ventilatory demand during exercise (21–24) (Figure 3). The rate and magnitude of DH during exercise is generally measured in the laboratory setting by serial inspiratory capacity (IC) measurements (3, 14, 25). Since total lung capacity (TLC) does not change during activity (23, 26), the change (decrease) in IC reflects the change (increase) in dynamic EELV, or the extent of DH. This simple method has been shown to be reliable and recent multi-centre clinical trials have confirmed its reproducibility and responsiveness (8, 14, 25). The use of change in IC to track DH is further validated by studies that have used esophageal manometry to demonstrate that even severely dyspneic patients are capable of generating maximal inspiratory pressures at the end of
Figure 2. Pressure-volume (P-V) relationships of the total respiratory system in health and in COPD. Tidal pressure-volume curves during rest (filled area) and exercise (open area) are shown. In COPD, because of resting and dynamic hyperinflation (a further increased EELV), exercise tidal volume (VT) encroaches on the upper, alinear extreme of the respiratory system's P-V curve where there is increased elastic loading. In COPD, the ability to further expand VT is reduced, i.e., inspiratory reserve volume (IRV) is diminished. In contrast to health, the combined recoil pressure of the lungs and chest wall in hyperinflated patients with COPD is inwardly directed during both rest and exercise; this results in an inspiratory threshold load on the inspiratory muscles. **Abbreviations:** EELV = end-expiratory lung volume; RV = residual volume; TLC = total lung capacity. Reprinted from Mahler DA, O'Donnell DE (eds). Dyspnea: Mechanisms, Measurement, and Management, 2nd edition. Lung Biology in Health and Disease Series, Volume 208, Chapter 3. New York: Taylor & Francis Group, 2005; pp. 29–58, used with permission.

exhaustive exercise (2, 27). This implies that the reductions in IC seen during exercise in COPD are not due to submaximal efforts, and indeed reflect changes in underlying EELV.

In combined studies conducted in over 500 patients with moderate-to-severe COPD, the change in EELV during cycle ergometry averaged 0.4 L, representing a reduction in IC by ~20% of the resting value, but with wide variation in the range (3, 5, 8) (Figure 4). Eighty-five percent of this population sample showed increases in EELV from rest to peak exercise, confirming the presence of significant DH (3, 5, 8). The minority of patients who showed little reduction in IC with exercise demonstrated the most severe resting lung hyperinflation (3). The rate of rise of DH was steeper in patients with the most severe expiratory flow-limitation (as estimated by the FEV₁/FVC ratio), the lowest diffusing capacity for carbon monoxide and the highest ventilatory demand (reflecting greater ventilation-perfusion abnormalities), and generally reached a maximal value early in exercise (3).

**Lung hyperinflation and activity limitation in COPD: The evidence**

The presence of expiratory flow-limitation appears to be an important predictor of exercise tolerance in patients with COPD. In a cohort of 52 patients, all of those who had evidence of significant expiratory flow-limitation at rest (measured by the negative expiratory pressure technique) showed a reduction of both peak workload and peak oxygen uptake (V'O₂) (28). By contrast, the achieved peak V'O₂ during exercise was within normal limits in 35% of the subjects who were not flow-limited at rest. Moreover, all of the patients with significant expiratory flow-limitation had evidence of resting lung hyperinflation: IC was reduced to <80% predicted. In contrast, IC was preserved in the majority of patients without flow-limitation. This supports the findings of Kouloris et al. (15) and suggests that: (1) reduced IC is a good and validated marker of flow-limitation and the propensity to develop worsening DH during exercise.
Changes in operating lung volumes are shown as ventilation increases with exercise in COPD (n = 105) and in age-matched normal subjects (n = 25). "Restrictive" constraints on tidal volume (V_t, solid area) expansion during exercise are significantly greater in the COPD group from both below (reduced inspiratory capacity (IC)) and above (minimal inspiratory reserve volume (IRV)). Other abbreviations: EELV = end-expiratory lung volume; EILV = end-inspiratory lung volume; Rrs = relaxation volume of the respiratory system; RV = residual volume; TLC = total lung capacity. With permission from O'Donnell DE, Webb KA. Mechanisms of dyspnea in COPD. In: Mahler DA, O'Donnell DE (eds). Dyspnea: Mechanisms, Measurement, and Management, 2nd edition. Lung Biology in Health and Disease Series, Volume 208, Chapter 3. New York: Taylor & Francis Group, 2005; pp. 29–58.

Figure 5. Significant intercorrelations between dyspnea intensity, neuromechanical dissociation and hyperinflation. Dyspnea, as assessed by the Borg scale, correlates significantly with the effort-displacement ratio (which is given by the ratio of Pes/PImax:VT/VC, where Pes is esophageal pressure, PImax is the maximal inspiratory pressure, VT is tidal volume, and VC is vital capacity) as an index of neuromechanical coupling. Dyspnea intensity is also significantly predicted by hyperinflation, as assessed by the end-expiratory lung volume (EELV) as a proportion of total lung capacity (TLC). The effort-displacement ratio and hyperinflation are also strongly correlated. From O’Donnell DE, Bertley JC, Chau LK, Webb KA. Am J Respir Crit Care Med 1997; 155:109–115 and adapted from Mahler DA, O’Donnell DE (eds). Dyspnea: Mechanisms, Measurement, and Management, 2nd edition. Lung Biology in Health and Disease Series, Volume 208, Chapter 3. New York: Taylor & Francis Group, 2005; pp. 29–58, used with permission.

Negative effects of acute dynamic hyperinflation during exercise

Although DH serves to optimise expiratory flow rates by avoiding expiratory flow-limitation at lower lung volumes, it has the deleterious effect of forcing VT to operate on the upper, flatter part of the respiratory system’s compliance curve where increases in pressure no longer generate significant incremental volume change (Figure 2). With worsening DH, the ability of the VT to increase during exercise is reduced, imposing “restrictive” mechanics. In fact, in 105 patients with COPD, the end inspiratory lung volume (EILV) was found to be 94 ± 5% of TLC at a peak symptom-limited VO2 of only 12.6 ± 5.0 mL/kg/min; this corresponded to a reduction in the inspiratory reserve volume (IRV) to a minimum of 0.42 ± 0.33 L (3). In contrast, when breathing at a minute ventilation (VE) similar to that of COPD patients at peak exercise, healthy age-matched individuals had significantly less constraints imposed on VT expansion, with IRV measured at 1.75 ± 1.16 L (Figure 3). DH reduces the ability of VT to expand appropriately during exercise and this leads to early mechanical limitation of VE (16). The consequence of this “saturation” of VT is that further increases in VE must rely on increases in breathing frequency (30). Unfortunately, in these already flow-limited patients, increases in breathing frequency may further aggravate DH in a vicious cycle and, in addition, contribute to reduced dynamic lung compliance and increased flow-resistive work.

DH results in sudden increases in the elastic and threshold loads on the inspiratory muscles, thus increasing the work and oxygen cost of breathing. The inspiratory threshold load (ITL) reflects the force that the inspiratory muscles must generate to counterbalance the inward (expiratory) recoil of the lung and chest wall at end-expiration and can be substantial in COPD (2). DH results in functional inspiratory muscle weakness by maximally shortening the muscle fibers in the diaphragm (31). The combination of excessive mechanical loading and increased velocity of shortening of the inspiratory muscles can also predispose them to fatigue. In some patients, this mechanical constraint on VT expansion, in the setting of severe ventilation-perfusion abnormalities (i.e., high fixed physiological dead space), leads to carbon dioxide retention and arterial oxygen desaturation during exercise (16). Finally, DH adversely affects dynamic cardiac function by contributing to pulmonary hypertension, by reducing right ventricular pre-load (reduced venous return) and, in some cases, by increasing left ventricular afterload (32–34). It has recently been postulated that competition between the over-worked ventilatory muscles with the active peripheral muscles.
for a reduced cardiac output may compromise blood flow and oxygen delivery to the latter, with negative consequences for exercise performance (35–37). All the above factors are clearly interdependent and contribute in a complex, integrated manner to dyspnea and exercise limitation in COPD.

Lung hyperinflation and exertional dyspnea in COPD: The evidence

Dyspnea is a complex multifaceted and highly personalized sensory experience, the source and mechanisms of which are incompletely understood. The notion that DH contributes to perceived exertional dyspnea has been bolstered by a number of studies that have shown a consistent statistical association between dyspnea intensity (assessed by the Borg scale) and various indices of DH during exercise (2, 3, 14). Using multiple regression analysis, subjective Borg ratings of dyspnea intensity were found to be strongly correlated with changes in EELV (expressed as % TLC; r = 0.63, p= 0.001) during exercise in 23 patients with advanced COPD (average FEV1 = 36% predicted). Furthermore, the measured change in EELV, and the subsequent constraint of VT expansion, also emerged as independent significant contributors to exertional dyspnea in these patients (1). In another study (2), exertional Borg dyspnea ratings correlated well with the ratio of EELV to TLC (r = 0.69, p < 0.001) (Figure 5). Similarly, in a larger study of 105 patients with moderate-to-severe COPD, the VT/IC ratio, as an index of VT constraint, emerged as the strongest predictor of exertional dyspnea (p < 0.0005) (3). Less important contributing variables included VfE/MVC, breathing frequency, and IRV/pred TLC, each accounting for 25% of the variance in Borg dyspnea ratings (p < 0.0005) (3). Poor correlation has also repeatedly been found between the FEV1 and/or the FVC and measures of disability such as dyspnea and exercise capacity (3).

Qualitative aspects of exertional dyspnea in COPD

Further insights into the link between dyspnea and DH in COPD have arisen from studies that have explored the qualitative aspects of respiratory discomfort at a point where it reaches intolerable levels at the end of exercise. In qualitative terms, patients with COPD consistently select descriptor clusters that allude to both “increased effort” and “unsatisfied inspiration” at the break-point of cycle exercise (Figure 6).

Dyspnea and perceived increased respiratory effort

Recent theories on the mechanisms of dyspnea have emphasized the central importance of the perception of increased contractile inspiratory muscle effort (38–45). When skeletal muscles are mechanically loaded, weakened or fatigued, increased electrical activation of the muscle is required to generate a given force, and motor output to these muscles is amplified. It is hypothesized that increased motor output is accompanied by increased corollary discharge to the sensory cortex where it is directly perceived as a heightened sense of effort (43–47). In COPD, inspired effort and central motor command output are increased compared with health, reflecting the relatively higher ventilation, the increased loading and functional weakness of the inspiratory muscles. Altered afferent information from activated mechanoreceptors in the overworked and shortened inspiratory muscles (secondary to DH) in COPD may contribute to an increased sense of work or effort, but this remains conjectural (47). Beyond a certain threshold, increased effort may be consciously registered as respiratory discomfort (38–43). Qualitative descriptors at end-exercise that allude to increased effort or work of breathing are pervasive across health and disease and increased corollary discharge remains a plausible mechanistic explanation for this (2) (Figure 6).

However, it must be remembered that increased sense of effort is only one component of this multi-dimensional symptom, and it is acknowledged that dyspnea can rise to severe levels even in the absence of increases in contractile muscle effort (48–54). Mechanical ventilation, which successfully unloads the ventilatory muscles (thereby reducing effort), may not fully alleviate dyspnea (55, 56). Chemoreceptor stimulation (by adding carbon dioxide) can induce breathing discomfort, described as air hunger, even in the absence of ventilatory muscle activation. Finally, increasing breathing effort to a high fraction of the maximal possible effort is not necessarily perceived as discomfort in all circumstances.

 Unsatisfied inspiration

In many respects, the sensory experience in COPD differs fundamentally from that of age-matched healthy individuals at peak VO2 (2). While the sense of increased effort, work or heaviness of breathing is common to both groups, only COPD patients consistently select descriptors that allude to unsatisfied inspiration (i.e., “can’t get enough air in”), and it is reasonable to assume that these different qualitative dimensions of exertional dyspnea in COPD reflect different underlying mechanisms (2).

The physiological events that occur at the end of exercise, when dyspnea becomes intolerable, are well understood. The neural drive to breathe reaches near maximal values, driven by the elevated carbon dioxide production (VCO2) that accompanies exercise and the early metabolic acidosis that may occur in many deconditioned COPD patients.

In some patients, critical arterial oxygen desaturation, sympathetic nervous system over-activation and altered feedback from peripheral muscle metaboreceptors may additionally stimulate ventilation. As already outlined, however, the ventilatory output in response to the increased drive is often markedly diminished because of derangements of dynamic ventilatory mechanics. It is noteworthy that, in contrast to health, the effort-displacement ratio (the ratio of inspired effort (tidal esophageal pressure relative to maximum inspiratory pressure, i.e., Pes/PImax) to volume displacement (tidal volume expressed as a percentage of predicted vital capacity, i.e., VT/VC)) continues to rise in COPD as exercise proceeds. This increased ratio, which crudely reflects the position of the operating tidal volume on the respiratory control system of breathing.
system’s pressure-volume relation (and thus the degree of neuromechanical dissociation), correlates well with perceived intensity of inspiratory difficulty. For example, in 12 patients with severe COPD (FEV\textsubscript{1} = 37% predicted), the effort-displacement ratio was the strongest correlate of dyspnea intensity during exercise (r = 0.86, p < 0.001), and also correlated strongly with dynamic hyperinflation (EELV/TLC; r = 0.78, p < 0.001) (2).

A recent mechanistic study in our laboratory has attempted to reconcile the beneficial effects of DH in early exercise with its deleterious sensory effects that ultimately contribute to exercise limitation. Thus, DH early in exercise allowed flow-limited patients to increase V\textsubscript{E} while minimizing respiratory discomfort (30). As a result of this early DH, the airways are maximally stretched at the higher lung volumes (close to TLC) and expiratory flow-limitation is attenuated allowing patients to maximize expiratory flow rates. Thus, patients with severe COPD could abruptly increase V\textsubscript{E} commensurate with increased metabolic demand, to approximately 40 L/min and generate tidal inspiratory pressures exceeding 40% of the maximal possible pressure generation while experiencing minimal increases in dyspnea (modified Borg ratings 1–2).

Effort-displacement ratios are therefore well maintained early in exercise even in advanced COPD. However, this advantage of DH was quickly negated when V\textsubscript{T} expanded to reach a critically low IRV of approximately 0.5 L (or 10% predicted TLC) below TLC (Figure 7). At this “threshold,” V\textsubscript{T} becomes fixed on the upper less compliant extreme of the respiratory system’s sigmoid-shaped pressure-volume relation, where there is increased elastic loading of the inspiratory muscles. At this operating volume, the diaphragm muscle fibers are maximally shortened and the increased breathing frequency leads to increased velocity of shortening and significant reductions in dynamic lung compliance. After reaching this minimal IRV, dyspnea (described as unsatisfied inspiration) soon rose to intolerable levels and reflected the widening disparity between inspiratory effort (reaching near maximal central neural drive) and the simultaneous V\textsubscript{T} response, which becomes essentially fixed, i.e., increased effort-displacement ratio (30). Consistent with a previous study (2), dyspnea intensity again correlated well with the increase in this effort-displacement ratio during exercise in COPD (30).

The contention that DH contributes importantly to exercise limitation in COPD has also been bolstered by numerous studies that have shown that pharmacological and surgical lung volume reduction are associated with consistent improvements in dyspnea and exercise endurance (4, 8, 14, 30, 57–63). Furthermore, reduced dyspnea ratings following bronchodilator therapy (i.e., tiotropium) were associated with consistent improvements in the effort-displacement ratio throughout exercise (30) (Figure 8).
Neurophysiology of exertional dyspnea

In health, during resting spontaneous breathing and during exercise, the mechanical output of the respiratory system, measured as $V_E$, changes in accordance with the level of central neural drive. Complex proprioceptive information (obtained from muscle spindles, Golgi tendon organs, and joint receptors), as well as sensory information pertaining to respired airflows and volume displacement (from mechanosensors located in the lung parenchyma and airways), provide simultaneous feedback to the central nervous system that ventilatory output is appropriate for the prevailing drive (46, 47, 64–68). Physiological adaptations during exercise, which include precise control of operating lung volumes and airway (intra- and extra-thoracic) resistance together with breathing pattern adjustments, ensure harmonious neuromechanical coupling of the respiratory system and avoidance of respiratory discomfort (69–71). Effort-displacement ratios therefore remain remarkably constant throughout exercise in health. Although the perceived effort of breathing may increase as $V_E$ increases during exercise, medullary output remains appropriately rewarded, and participants generally do not describe inspiratory difficulty or unsatisfied respiratory effort, even at peak exercise (2).

The situation is markedly different in COPD, where DH during exercise constrains $V_T$ expansion and results in maximal shortening of the inspiratory muscles. Once $V_T$ expands to reach a critical IRV ceiling, further increases in neural output to the respiratory system are unrewarded in terms of increased mechanical output. We have argued that this mechanical (volume) restriction is a primary mechanism by which DH induces exertional dyspnea and its dominant qualitative dimension of unsatisfied inspiration (2, 30, 72). It is possible, therefore, that sensory feedback from a multitude of mechanoreceptors throughout the respiratory system (in the muscles, chest wall, airways and lung parenchyma) collectively convey the information to consciousness that the mechanical output achieved is inadequate for the prevailing respiratory drive. In the final phase of exercise, central drive had likely reached near maximal levels yet the $V_T$ response was essentially fixed at only 30% of the predicted vital capacity (30). Respiratory mechanoreceptors are ideally placed to detect any disparity between the volume displacement achieved and that which is expected (73).

Dyspnea and volume restriction in COPD

Several previous studies in resting healthy humans have shown that when chemical drive is increased in the face of voluntary suppression or imposed restriction of the spontaneous breathing response (i.e., $V_T$ expansion), dyspnea quickly escalates to intolerable levels (48–50, 72, 74, 75). Moreover,
resumption of spontaneous breathing was associated with immediate improvement in respiratory discomfort, despite persistent (or even increased) chemical loading. During exercise in health, mechanical restriction of VT (by chest strapping) induced severe dyspnea (described as unsatisfied inspiration) in the setting of added chemical loading (72). We postulate that in COPD, a similar mismatch between central drive and a restricted mechanical response (as a result of DH) is fundamental to the origin of dyspnea. This hypothesis is supported by a number of controlled therapeutic studies that have shown a correlation between reduced dyspnea intensity ratings and the extent of release of tidal volume restriction following pharmacological lung volume reduction (4, 8, 14, 30, 57–60).

It is certainly possible that acute DH could result in fatigued or excessively shortened inspiratory muscle fibers that fail to respond appropriately to increased electrical activation. Accordingly, as originally proposed by Campbell (76), spindles in the ventilatory muscles (which accurately sense the disparity between length and tension development) are ideally suited to serve as the proximate peripheral source of this sensory information (42, 46, 47, 70, 71, 73, 74). However, based on the existing literature, it is not clear whether overt inspiratory muscle fatigue consistently occurs in the setting of symptom-limited exercise, even in severe COPD (77–80). Moreover, it is highly unlikely, given the considerable redundancy in the neurosensory system, that muscle spindles are the only mechanosensors that provide sensory feedback with respect to the appropriateness of the ventilatory output for the prevailing drive.

**Dyspnea: The emotional dimension**

We have seen that DH-associated dyspnea is likely to arise in flow-limited patients under conditions of abruptly increased neural drive and ventilatory demand. Common provocative situations include: physical activity, episodes of transient hypoxemia and anxiety. Recent studies have confirmed that daily fluctuations in DH can reflect circadian variability in airway smooth muscle tone that is not fully reversed by long-acting bronchodilators. More sustained DH occurs in the setting of exacerbations and this has recently been linked to dyspnea, the dominant presenting symptom (18, 19). It is reasonable to assume that when perceived respiratory discomfort exceeds a certain threshold (which varies between individuals), it will elicit behavioral or affective responses. This affective dimension, in many instances encompasses feelings of fear that can quickly escalate to panic and helplessness, which are key components of perceived respiratory distress. Sudden fear or overt panic will elicit neuro-humoral responses (via pathways in the amygdala, adrenals and sympathetic nervous system), which will trigger patterned ventilatory and circulatory responses that can further amplify respiratory discomfort.

Recently, the use of functional imaging techniques such as positron-emission tomography (PET) scanning and functional magnetic resonance imaging (fMRI) have been utilized to investigate the mechanisms underlying the central processing and perception of dyspnea (81–86). These studies have shown activation of central limbic structures including the anterior insula, pars opercularis, anterior cingulate gyrus, amygdala, putamen, and caudate. These phylogenetically ancient areas of the central nervous system have an integral role in the perception and genesis of primal emotions, and it has been suggested that air hunger and dyspnea evoke programmed neurohumoral and behavioral responses similar to those that occur in response to pain (87–89), extreme hunger (90) or thirst (91). Other data suggest that the anterior insula is also activated in the setting of panic attacks (92), which may provide a common pathway for the disabling sensations of panic, anxiety and fear that often accompany severe dyspnea (93).

**SUMMARY**

Severe dyspnea is a major exercise-limiting symptom in moderate-to-severe COPD and every effort should be made to alleviate this distressing symptom. Although exercise limitation is multifactorial, there is considerable evidence that deranged ventilatory mechanics, specifically dynamic lung hyperinflation, may represent the proximate mechanical limit to exercise performance in patients with more advanced disease. Dynamic lung hyperinflation occurs during activity in the vast majority of flow-limited patients with COPD and has been shown repeatedly to correlate with dyspnea intensity ratings. Dynamic lung hyperinflation stresses the already limited cardiopulmonary reserves.
of patients with COPD and greatly constrains their ability to expand tidal volume appropriately in response to the increased neural drive of exercise. Recent studies have proposed that this acute neuromechanical dissociation of the respiratory system may form the basis for the perception of respiratory discomfort, which ultimately triggers intolerable respiratory distress. Dynamic lung hyperinflation therefore represents an important therapeutic target in COPD and several studies have now shown that pharmacological lung deflation is associated with clinically important improvements of dyspnea and exercise endurance, even in advanced disease.

REFERENCES
33. Light RW, Mintz WM, Linden GS, Brown SE. Hemodynamics of patients with severe chronic obstructive pulmonary disease during


