Use of cluster analysis to describe desaturator phenotypes in COPD: correlations between pulmonary function tests and nocturnal oxygen desaturation

Domenico Maurizio Toraldo1
Francesco De Nuccio2
Annarita Gaballo1
Giuseppe Nicolardi2

1A Galateo Lung Disease Hospital, Regional Service Puglia, San Cesario di Lecce; 2Laboratory of Human Anatomy, Department of Biological and Environmental Sciences and Technologies, University of Lecce, Lecce, Italy

Background: Significant heterogeneity of clinical presentation and disease progression exists within chronic obstructive pulmonary disease (COPD). Although forced expiratory volume in 1 second (FEV1) inadequately describes this heterogeneity, a clear alternative has not emerged. This article discusses and refines the concept of phenotyping desaturators in COPD and shows a possible pattern which could be used as a framework for future research.

Recent findings: COPD is a complex condition with pulmonary and extrapulmonary manifestations. We suggest that COPD phenotypes should be associated with clinically meaningful outcomes. The innovation of COPD phenotyping is defined as COPD desaturators. Sleep-related hypoxemia and hypercapnia are well recognized in COPD and the development of systemic inflammation during sleep. These sleep-related changes predispose to nocturnal cardiac arrhythmias, pulmonary hypertension, and possibly death, particularly during acute exacerbations.

Conclusion: A more focused definition makes possible a classification of patients into two distinct subgroups for both clinical and research purposes. Establishing a common language for future research will facilitate our understanding and management of such diseases. Even if different treatment strategies have different outcomes for these groups, we will have confirmation, or otherwise, of the clinical value of cluster analysis. This knowledge could lead to pharmacological treatment and other interventions directed to specific phenotypic groups.

Keywords: phenotypes, chronic obstructive pulmonary disease, desaturator, nocturnal hypoxemia, systemic inflammation, intermittent hypoxia

Introduction

Chronic obstructive pulmonary disease (COPD) is defined as a “preventable” but treatable disease with significant extrapulmonary effects that may contribute to its severity in individual patients.1 However, COPD is a complex, multicomponent, heterogeneous disease, the clinical, functional and radiological presentation of which varies greatly from patient to patient, even though the degree of airflow limitation may be similar.1,2 The current classification of airways disorders is imprecise, with an overlap of phenotypes (eg, asthma, chronic bronchitis, and emphysema), resulting in difficulties in differentiating between these disorders.

The prevalence, distribution, and interrelationships of the main clinical and functional manifestations of the disease in a large, well-characterized, and controlled population of patients is lacking. Comorbidities such as chronic heart failure, cardiovascular disease, depression, diabetes, muscle wasting, weight loss, lung cancer, and...
Systemic inflammation in COPD

Several studies have found markers of systemic inflammation, such as high-sensitivity C-reactive protein, to be higher in the blood of patients with COPD than in subjects without COPD. The question arises as to whether systemic inflammation is the result of a local inflammation spillover into the systemic compartments or a systemic component of COPD not necessarily related to local inflammatory processes in the lung. As a point of interest, it should be noted that systemic inflammation has failed, so far, to show substantial correlations with airway obstruction, whereas at least some relationship has been reported between local inflammatory processes and airway obstruction. Markers of systemic inflammation have been shown to be elevated in the blood of patients with COPD when compared with control subjects without COPD. COPD is often accompanied by other chronic diseases associated with systemic inflammation, such as chronic heart failure, diabetes, and arteriosclerosis. Alternatively, increased blood levels of inflammatory mediators in patients may stem from extrapulmonary cells (circulating leukocytes, endothelium, or muscle cells). A particular problem in COPD patients with marked alveolar wall destruction is intermittent and continuous hypoxia. A significant inverse correlation between arterial oxygen tension (PaO₂), circulating tumor necrosis factor alpha, and soluble tumor necrosis factor receptor levels has been reported in patients with COPD. Similarly, a significant relationship between reduced oxygen delivery and tumor necrosis factor alpha levels in the peripheral circulation has been found, highlighting the role of nocturnal hypoxia in the tissue. It has been suggested that systemic inflammation may explain part of the heterogeneity of COPD phenotypes, such as loss of lean body mass and the higher prevalence of comorbid disorders, such as coronary heart disease, depression, and hypertension. Finally, this study reinforces the view that systemic inflammation is an important phenotypic feature of COPD. Future prospective studies should investigate whether these markers give important prognostic information in relation to disease progression and severity in COPD.

Sleep in patients with COPD

COPD alone can cause subjective and objective changes during sleep. When patients with chronic bronchitis or emphysema were surveyed across a broad range of symptoms, “sleep difficulties” were endorsed as occurring “almost always” or “always” in 43% of subjects (third most common, after dyspnea and fatigue). In addition to the diagnosis of COPD, the presence of COPD symptoms, such as cough, sputum production, and wheezing correlated strongly with difficulty in falling or staying asleep. Other investigations have objectively confirmed poor sleep quality, with decreased total sleep time and decreased sleep efficiency.

A brief review of the normal changes in respiration that occur with sleep onset and the various sleep stages is useful for understanding the changes occurring during sleep in patients with COPD. In healthy subjects, minute ventilation drops from wakefulness to non-rapid eye movement (non-REM) sleep, and drops further during REM sleep (about 15% compared with the awake value). Most of the drop in minute ventilation is due to a decrease in tidal volume not fully compensated by a concomitant increase in respiratory rate. There is a blunted ventilatory response to hypoxia and hypercapnia, again with the greatest changes during REM sleep. Sleep-related hypoventilation has been demonstrated in COPD, particularly during REM sleep, with associated oxygen desaturation. Nocturnal oxygen desaturation in COPD is likely to be the consequence of the combined effects of physiological hypoventilation during sleep. However, there is evidence that some patients with awake PaO₂ levels in the mildly hypoxemic range can also develop clinically significant nocturnal oxygen desaturation, which may predispose to pulmonary hypertension. Finally, possible mechanisms responsible for this reduction include respiratory muscle hypotonia, cephalic displacement of the diaphragm, and a decrease in lung compliance.
Mechanisms of sleep-related breathing disturbances in COPD

Sleep-related hypoventilation has been demonstrated in COPD, with associated oxygen desaturation particularly during REM sleep. There is a close relationship between the awake PaO₂ and nocturnal oxygen saturation levels, although hypercapnia is associated with more pronounced nocturnal oxygen desaturation than normocapnia for any given level of waking oxygen saturation level.

Nocturnal hypoxemia is defined as an oxygen saturation level ≤90% for at least 5 minutes, with a nadir oxygen saturation level of ≤85%. Time in bed is defined as the time from the start to the end of the recording. The percentage of total recording time is defined as time spent in bed – sleep latency + intrasleep wakefulness. The total recording time spent in bed with an oxygen saturation level ≤90% is defined as desaturators and the others as nondesaturators. COPD desaturator patients may be identified by a clinical pattern of variables rather than by T₉₀ alone, ie, T₉₀ mean pulmonary artery pressure, and arterial carbon dioxide tension values, with the latter two variables being predictors of severity of nocturnal desaturation. Alveolar hypoventilation probably accounts for most of the oxygen desaturation. Becker et al measured minute ventilation during wakefulness, non-REM sleep, and REM sleep in normal subjects and in patients with COPD. The greater drop seen in minute ventilation in subjects with COPD may reflect increased dependence on accessory muscles that become hypotonic during sleep, particularly during REM sleep. An alternative explanation comes from the work of O’Donoghue et al who found an even greater drop in minute ventilation during non-REM sleep in hypercapnic COPD patients.

Consequences of nocturnal oxygen desaturation in COPD

The exact prevalence of pulmonary hypertension in patients with COPD is unclear. Pulmonary hypertension is a complication of advanced COPD observed in patients who show severe longstanding hypoxemia. Even if pulmonary hypertension is generally mild to moderate in most COPD patients, it may worsen markedly during acute exacerbations, sleep, and exercise, and these acute increases in pulmonary hypertension could facilitate the development of right heart failure. Diagnosis of pulmonary hypertension in COPD patients is difficult. The published studies differ not only in their definitions but also for conditions in which pulmonary hypertension has been reported (ie, rest, exercise, and exacerbation). According to the European Society Cardiology and European Respiratory Society, pulmonary hypertension is defined as an increase in mean pulmonary artery pressure ≥25 mmHg at rest as assessed by right heart catheterization. The definition of pulmonary hypertension on exercise as a pulmonary artery pressure ≥30 mmHg assessed by right heart catheterization is not supported by the published data, and healthy individuals can reach much higher values.

The incidence of pulmonary hypertension in COPD patients has been evaluated by Kessler et al who performed a longitudinal study in 131 patients with COPD with serial right heart catheterization at baseline and then at follow-up (mean 6.8 ± 2.9 years). All subjects had normal mean pulmonary artery pressure at rest (≤20 mmHg). They were divided into two groups according to the presence or absence of elevated mean pulmonary artery pressure with exercise (≥30 mmHg), and 25% of patients developed pulmonary hypertension on follow-up that was mild by hemodynamic criteria (mean pulmonary artery pressure 26.8 ± 6.6 mmHg). Subjects who showed elevated pulmonary hypertension with exercise were more likely to exhibit resting mean pulmonary artery pressure elevation at follow-up. The annual rate of progression was ±0.4 mmHg.

Nocturnal oxygen desaturation seems to contribute to the development of pulmonary hypertension, even in the absence of significant awake hypoxemia. REM-associated falls in oxygen saturation levels are associated with increases in pulmonary artery pressure during sleep that can be reversed by supplemental oxygen, although most COPD patients with sustained pulmonary hypertension are also hypoxemic during the daytime. Various arrhythmias have also been reported during episodes of nocturnal desaturation. These observations might help to explain why nocturnal oxygen desaturation is a marker of increased mortality, and why COPD patients are reported to die more frequently than expected at night.

Tissue hypoxia is another mechanism that can contribute to systemic inflammation in COPD. In one clinical study it was shown that tumor necrosis factor alpha and its receptor levels were significantly higher in patients with COPD, and were correlated significantly with severity of arterial hypoxemia. These results suggest that arterial hypoxemia in COPD is associated with activation of the tumor necrosis factor alpha system in vivo.
The systemic effects of inflammation may contribute significantly to not only the respiratory abnormalities, symptoms, and functional impairment (eg, exercise intolerance) associated with COPD but also to the marked changes in vasomotor and endothelial function seen in chronic pulmonary vascular disease. The nocturnal desaturation-reoxygenation sequence is a typical pattern coupled with the majority of respiratory events. This sequence, defining intermittent hypoxia, leads to oxidative stress, with production of reactive oxygen species. Hypoxia-induced pulmonary vasoconstriction is a protective response to keep the ventilation-perfusion ratio at an optimum level by shunting blood away from hypoxemic areas. The traditional hypoxic model of pulmonary hypertension is based on the hypothesis that chronic hypoxia initiates vascular remodeling, leading to permanent changes in the pulmonary vasculature. Studies performed in vitro have elucidated the mechanisms underlying hypoxia-driven vascular changes. Barbera et al evaluated COPD patients undergoing lung resection and demonstrated that vascular changes contribute to vascular remodeling and putatively may have an effect on the vascular dynamics leading to pulmonary hypertension. Nocturnal hypoxia may induce endothelial cells to release cytokines, leading to cellular hypertrophy in the vessel wall and an increase in the extracellular matrix. In conclusion, the nocturnal hypoxic insult occurring during sleep-disordered breathing may contribute to chronic vascular remodeling, causing vascular endothelial damage and dysfunction, and increasing the risk of pulmonary hypertension in COPD.

**Use of cluster analysis to define potential COPD phenotypes**

Traditionally, on the basis of specific clinical, functional, and radiological features, patients with COPD used to be classified into different biotypes, ie, the “blue bloater,” in association with predominantly chronic bronchitis, and the “pink puffer,” identified as predominantly emphysema. However, these are only two extreme phenotypes among a broader spectrum of clinical presentations in COPD. Accordingly, we propose a variation on the traditional definition of a phenotype, ie, a single disease attribute or a combination of disease attributes describing differences between individuals with COPD as they relate to clinically meaningful outcomes. It is proposed that phenotypes in COPD should have real predictive value. Studies carried out in recent years have revealed that patients with the same stage of disease may show different pathological changes, and classic COPD phenotypes clearly differ from these based on severity of emphysema as assessed by high-resolution computed tomography scanning.

The goal of phenotyping is to identify patient groups with unique prognostic or therapeutic characteristics. However, significant variation and confusion surrounds the use of the term “phenotype” in COPD. Phenotype classically refers to any observable characteristic of an organism, and until now multiple disease characteristics have been termed COPD phenotypes. Evidence has shown that different COPD phenotypes may be significantly associated with differences in body mass index, health-related quality of life, small airways obstruction, and systemic inflammation.

Cluster analysis is a collection of methods for defining groups of individuals based on measured characteristics, so that they can be grouped according to their similarities or differences into clusters. Groupings are created so that the degree of association is strong between members of the same cluster and weak between members of different clusters. Cluster analysis is distinct from other ways of trying to understand multivariate data, including principal component and factor analysis, discriminant analysis, and multivariate regression. The clinical relevance of cluster analysis will depend on developing diagnostic criteria to allow new individuals to be allocated into groups based on the identified clusters.

More recent work, including that of Burgel et al, has used cluster analysis to characterize different types of airways disorder. The main conclusion from Burgel et al is that COPD patients with similar airflow obstruction can belong to different phenotypes, and have different symptoms (dyspnea) and outcomes (number of exacerbations and predicted mortality). At a more specific level, it is worthy of note that both Wardlaw et al and Weatherall et al have identified a cluster characterized by severe and markedly variable airflow obstruction with features of atopic asthma, chronic bronchitis, and emphysema. Patients in this phenotypic group would be unlikely to meet the inclusion criteria of the major randomized, controlled trials of either asthma or COPD. The identification of COPD phenotypes will require an iterative validation process in which candidate phenotypes are identified before their relevance to clinical outcome is determined.

**Classification of COPD desaturator phenotypes using cluster analysis**

COPD patients can be classified as desaturators or non-desaturators based on a T90 of 30% on polygraphic recording.
One study\textsuperscript{29} of 51 consecutive COPD outpatients with mild daytime hypoxemia (PaO\textsubscript{2} 60–70 mmHg) identified a pattern of daytime clinical variables that distinguished desaturators from nondesaturators by using cluster analysis (see Table 1).

Considering all patients, T\textsubscript{90}, mean pulmonary artery pressure at rest, PaCO\textsubscript{2}, nadir oxygen saturation level, mean nocturnal oxygen saturation level, predicted total lung capacity, baseline oxygen saturation level awake, predicted vital capacity, and body mass index had a bimodal distribution. The variables examined in the study did not differ between men and women. Rather than using T\textsubscript{90} alone, desaturator patients may be identified by a pattern of T\textsubscript{90} (30.08\%–45.1\%, P = 0.0001) mean pulmonary artery pressure (33.1 ± 0.7 mmHg, P = 0.0006), and PaCO\textsubscript{2} (35.0–57.9 mmHg, P = 0.0005) values, with the latter two variables being predictors of severity of nocturnal desaturation (see Table 2).

A hierarchical cluster analysis has been performed using these three variables, showing the highest correlation by least-squares multiple linear regression (ie, T\textsubscript{90}, mean pulmonary artery pressure, and PaCO\textsubscript{2}). First, we identified two major clusters of patients designated as desaturator and nondesaturator, their mean values coinciding with those of the cluster centroid. In any event, a T\textsubscript{90} cutoff value does not appear to describe a cluster of desaturator patients adequately or to assess correctly the severity of nocturnal desaturation. Moreover, cluster analysis identifies subgroups of desaturator and nondesaturator patients who differed in degree of disease severity.

The finding that T\textsubscript{90}, mean pulmonary artery pressure, and PaCO\textsubscript{2} were required to identify desaturator and non-desaturator patients by cluster analysis demonstrates that these variables play a predictive role. Desaturator patients have higher values for mean pulmonary artery pressure, PaCO\textsubscript{2}, and T\textsubscript{90} than nondesaturator patients. Interestingly, cluster analysis has identified subpopulations of desaturator and nondesaturator patients, ie, two desaturator subgroups divided according to mean pulmonary artery pressure values, and two nondesaturator subgroups divided according to PaCO\textsubscript{2} values. By reducing the distance between the elements of clusters, more clusters (or rather subdivisions) of desaturator and nondesaturator groups were obtained.\textsuperscript{30}

Two subgroups, accounting for 76\% of patients, were identified in the nondesaturator group, with the remaining 24\% (six of 25 patients) being comprised of scattered individual patients. The ND1 nondesaturator subgroup comprised five of 25 patients (20\%; one of five women [20\% of the subgroup]; four of five men [80\% of the subgroup]). The centroids were as follows: T\textsubscript{90}, 22.0\%; mean pulmonary artery pressure, 19.2 mmHg; and PaCO\textsubscript{2}, 46.5 mmHg. The ND2 nondesaturator subgroup comprised 14 of 25 patients (56\%; eight of 14 women [57.1\% of the subgroup]; six of 14 men [42.9\% of the subgroup]). The centroids were as follows: T\textsubscript{90}, 20.1\%; mean pulmonary artery pressure, 18.5 mmHg; and PaCO\textsubscript{2}, 33.3 mmHg. Similarly, the desaturator group

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Variable mean values ± standard errors of the mean and range of both desaturator and nondesaturator groups of COPD patient phenotypes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Variables</td>
<td>Group D (n = 26)</td>
</tr>
<tr>
<td>Age (years)</td>
<td>Mean ± SD</td>
</tr>
<tr>
<td>70.5 ± 3.9</td>
<td>71.0</td>
</tr>
<tr>
<td>BMI kg/m(^2)</td>
<td>31.3 ± 2.1</td>
</tr>
<tr>
<td>PaO\textsubscript{2} mmHg</td>
<td>63.5 ± 3.4</td>
</tr>
<tr>
<td>PaCO\textsubscript{2} mmHg</td>
<td>50.0 ± 4.2</td>
</tr>
<tr>
<td>Baseline SaO\textsubscript{2} awake (%)</td>
<td>89.9 ± 1.9</td>
</tr>
<tr>
<td>FEV\textsubscript{1} (% predicted)</td>
<td>52.2 ± 9.6</td>
</tr>
<tr>
<td>FEV\textsubscript{1}/FVC ratio</td>
<td>40.1 ± 3.4</td>
</tr>
<tr>
<td>TLC (% predicted)</td>
<td>107.4 ± 7.9</td>
</tr>
<tr>
<td>Vital capacity (% predicted)</td>
<td>84.7 ± 6.8</td>
</tr>
<tr>
<td>MPAP at rest (mmHg)</td>
<td>33.1 ± 3.6</td>
</tr>
<tr>
<td>Mean nocturnal SaO\textsubscript{2} (%)</td>
<td>86.6 ± 4.1</td>
</tr>
<tr>
<td>Nadir SaO\textsubscript{2} (%)</td>
<td>78.4 ± 7.2</td>
</tr>
<tr>
<td>T\textsubscript{90} (%)</td>
<td>37.2 ± 3.5</td>
</tr>
<tr>
<td>AHI, per hour of sleep</td>
<td>2.9 ± 0.8</td>
</tr>
<tr>
<td>ESS score</td>
<td>3.9 ± 1.2</td>
</tr>
</tbody>
</table>

Abbreviations: AHI, apnea-hypopnea index; BMI, body mass index; ESS, Epworth Sleepiness Scale; FEV\textsubscript{1}, forced expiratory volume in 1 second; FVC, forced vital capacity; MPAP, mean pulmonary artery pressure; TLC, total lung capacity; T\textsubscript{90}, recording time spent in bed with an oxygen saturation level ≤90\%; PaO\textsubscript{2}, arterial oxygen tension; PaCO\textsubscript{2}, arterial carbon dioxide tension; SaO\textsubscript{2}, oxygen saturation; COPD, chronic obstructive pulmonary disease; D, desaturator; ND, nondesaturator; SD, standard deviation.
Table 2 Variables describing desaturator COPD patient phenotypes using cluster analysis

<table>
<thead>
<tr>
<th>Variables</th>
<th>Group D</th>
<th>Group ND</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Mean ± SE</td>
<td>Range</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean pulmonary artery pressure (PaCO2), mmHg</td>
<td>37.2 ± 0.7</td>
<td>30.8–45.5</td>
<td></td>
</tr>
<tr>
<td>MPAP at rest, mmHg</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean pulmonary artery pressure (PaCO2), mmHg</td>
<td>33.1 ± 0.7</td>
<td>28.0–40.0</td>
<td></td>
</tr>
<tr>
<td>PaCO2, mmHg</td>
<td>50.0 ± 0.8</td>
<td>35.1–57.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>21.2 ± 3.6</td>
<td>15.2–28.4</td>
<td>0.0001</td>
</tr>
<tr>
<td></td>
<td>19.8 ± 4.3</td>
<td>16.0–37.0</td>
<td>0.0006</td>
</tr>
<tr>
<td></td>
<td>38.0 ± 6.9</td>
<td>30.2–50.9</td>
<td>0.0005</td>
</tr>
</tbody>
</table>

Abbreviations: SE, standard error of the mean; MPAP, mean pulmonary artery pressure; PaCO2, arterial carbon dioxide tension; COPD, chronic obstructive pulmonary disease; D, desaturator; ND, nondesaturator.

was divided into two subgroups accounting for 88.4% of desaturator patients, with the remaining 11.5% (three of 26 patients) consisting of scattered individual patients. The D1 desaturator subgroup comprised seven of 26 patients (26.9%, three of seven women [42.8% of the subgroup]; four of seven men [57.2% of the subgroup]). The centroids were as follows: T90, 38.8%; mean pulmonary artery pressure, 38.0 mmHg; and PaCO2, 52.0 mmHg. The D2 desaturator subgroup comprised 16 of 26 patients (61.5%; five of 16 women [31.2% of the subgroup]; 11 of 16 men [68.8% of the subgroup]). The centroids were as follows: T90, 37.0%; mean pulmonary artery pressure, 31.5 mmHg; and PaCO2, 49.9 mmHg. These data show that PaCO2 was higher in ND1 patients than in ND2 patients, and that mean pulmonary artery pressure was higher in D1 patients than in D2 patients.

Finally, cluster analysis showed that most of the subjects with higher PaCO2 values were men in the small ND1 subgroup. This applies also to the D2 subgroup, but the men in this subgroup had lower levels of mean pulmonary artery pressure levels than the women.

Further studies are required to understand the importance of these findings. They provide a naturalistic classification that, if confirmed in other studies, could be developed into a modified taxonomy for disorders of sleep-related breathing disturbances in COPD. In conclusion,39 this cluster analysis showed that clustered COPD desaturator patients can be identified not by T90 value alone, but by a pattern of T90, mean pulmonary artery pressure, and PaCO2 values, and that the latter two variables are predictors of the severity of nocturnal desaturation.

Awake desaturator patients have lower PaO2 and higher PaCO2 values than awake nondesaturator patients.57 Moreover, daytime hypercapnia is a risk factor for nocturnal hypoxemia in COPD patients with mild daytime hypoxemia.58

The degree of airways obstruction in COPD patients, as measured by forced expiratory volume in 1 second (FEV1)/forced vital capacity, can correlate with the risk of prolonged hypoxemia,59 which appears to increase the morbidity and mortality risk in these patients, although the exact mechanism(s) that account for this increased risk are not well understood. The increased risk of death may be due to more prolonged hypoxia, although night-time hypercapnia is probably also greater. There is also increasing evidence that COPD has systemic consequences. Inflammation is caused via various mediators (tumor necrosis factor alpha, interleukin-6, and interleukin-8), in addition to the oxidative stress. The review by McNicholas shows how the disease acts through similar pathways to cause cardiovascular disease.60

Another intriguing possibility mentioned in that review is that nocturnal desaturation in COPD may contribute to an increased incidence of COPD exacerbations, which may accelerate the decline in lung function and be associated with greater mortality.61,62

Further cluster analyses, both population-based and clinic-based, will contribute to greater understanding of the true patterns of airway disorders. The clinical application of cluster analysis will depend on developing diagnostic criteria to allow new individuals to be allocated to groups based on identified clusters.

Diagnosis

The Global Initiative for Chronic Obstructive Lung Disease (GOLD)1 guidelines suggest that patients with relatively mild COPD and evidence of pulmonary hypertension and diurnal hypercapnia should be referred for overnight testing. This recommendation reflects data collected by Chaouat et al66 and Toraldo et al30 and emphasized by Kessler,35 ie, in all COPD patients with obesity, higher diurnal PaCO2 reflects increased muscle load in those with both increased upper and lower airway resistance. It is necessary to perform the following examinations:30 static lung volumes measured by body plethysmography and dynamic lung volumes by mass flow sensors in seated patients according to standard procedures; arterial blood gases measured at the radial artery using microelectrodes in seated patients spontaneously breathing air; resting mean daytime pulmonary artery pressure measured by color
Doppler echocardiography; and nocturnal desaturation evaluated by polygraphic recording of oxygen saturation, snoring, air flow, thoracic and abdominal respiratory movements, heart rate, including an electrocardiogram in real-time mode; and body position.

In our research, patients with a history of loud snoring and excessive daytime sleepiness, as evaluated by the Epworth Sleepiness Scale (range 0–10) and with an apnea-hypopnea index ≥5 per hour were excluded because of suspected obstructive sleep apnea syndrome. We also excluded patients in whom mean pulmonary artery pressure could not be evaluated by color Doppler echocardiography. Finally, quantitative assessment of emphysema by computed tomography scanning offers an objective measure of parenchymal disease that correlates well with histopathologic findings and is predictive of the degree of expiratory airflow obstruction. Objective measures of proximal airway wall thickening obtained via computed tomography are inversely correlated with lung function and relate to burden of small airway disease and exacerbation frequency.

**Treatment**

Management options for patients with sleep-related respiratory disturbances include general measures, such as optimizing therapy for the underlying condition and supplemental oxygen, in addition to pharmacological therapy. We propose that COPD phenotypes should be associated with clinically meaningful outcomes. Such a more focused definition makes possible a classification of patients into distinct prognostic and therapeutic subgroups for both clinical and research purposes. The goal of treatment is to maintain adequate oxygenation at all times and to prevent sleep-disordered breathing. Supplemental oxygen is the mainstay of treatment for those with daytime and nocturnal hypoxemia, and has been shown to reduce overall mortality if used for more than 18 hours per day, including during sleep.63,64

It may be that COPD patients with hypoxemia are at increased risk of mortality only during sleep, compared with those who are not, although this finding is only based on a single study of retrospective data.65 Again, correction of nocturnal hypoxemia alone (in patients with daytime normoxia) does not seem to improve pulmonary hemodynamics or mortality to a significant extent,66,67 although it may improve sleep quality and is frequently prescribed.68

A Cochrane analysis69 determined the effect of domiciliary oxygen on survival in COPD and identified six relevant randomized controlled trials. There was no effect on survival during 3 years of follow-up in patients with mild to moderate hypoxemia and those with only arterial desaturation at night. Home oxygen improved survival only in patients with \( \text{PaO}_2 \leq 60 \text{ mmHg} \).

Controversy exists as to whether continuous positive airway pressure (CPAP) therapy improves daytime lung function in patients with stable COPD. At least in an animal model, upper airways irritation increased lower airway resistance, so, in theory, correction of repetitive airway collapse might improve pulmonary function.70 Others have postulated that offloading the respiratory muscles could decrease hypoventilation, oxygen consumption, and carbon dioxide production by the respiratory muscles. These muscles may be rested by CPAP, given that it prevents the increase in upper airway resistance occurring during sleep. Alternatively, CPAP may offset intrinsic positive end-expiratory pressure in severe COPD. Mezzanotte et al used CPAP for 1–3 weeks in eight patients with COPD and assessed inspiratory force and endurance. They found significant improvements in maximum inspiratory force and performance on a 12-minute walk test.71 Improvements have also been observed in daytime oxygenation and hypercapnia.72,73

Long-term nocturnal noninvasive ventilation can also be considered in COPD patients with chronic respiratory failure, with improvements in gas exchange during wakefulness having been reported,74 in addition to improvements in respiratory muscle strength and endurance.75 Sleep quality and diurnal \( \text{PaO}_2 \) and \( \text{PaCO}_2 \) levels are better with noninvasive ventilation plus supplemental oxygen than with supplemental oxygen alone.76 There has been considerable interest in the use of noninvasive ventilation in stable hypercapnic COPD, with multiple studies and inconsistent results over the years.77 Several mechanisms are likely to play a role in these improvements, including rest of chronically fatigued respiratory muscles, thereby improving daytime respiratory muscle function.78

Two recent studies warrant attention. The first was a randomized controlled trial of noninvasive ventilation in patients with stable hypercapnic COPD by McEvoy et al, which showed significant improvement in adjusted mortality.79 There was little or no change in pulmonary function or daytime blood gases. The improvement in mortality using noninvasive ventilation was associated with a worse quality of life, which tempers enthusiasm for this approach. A second report by Windisch et al reported a reduction in mortality with noninvasive ventilation, although only used historical controls. However, these authors used what they called “high-intensity noninvasive ventilation,” with a very high driving pressure (average inspiratory pressure 28 cm \( \text{H}_2\text{O} \), average expiratory pressure 5 cm \( \text{H}_2\text{O} \)) and a high respiratory rate (about 21 breaths per minute). With
those settings, requiring hospital acclimatization, there were improvements in spirometry and blood gas abnormalities.\textsuperscript{80}

**Weight loss**

Weight loss can clearly be beneficial in obese COPD patients with a desaturator phenotype.\textsuperscript{81} However, in COPD, weight loss has generally been associated with increased mortality, because cachexia sets in with increasing disease severity. Thus, there are no data to recommend weight loss as a therapeutic option in those with the COPD desaturator phenotype, although it seems reasonable that those with less severe COPD would benefit from dietary and respiratory rehabilitation.

**Bronchodilators and corticosteroids**

Treatment of the underlying obstructive lung disease is helpful in preventing or ameliorating nocturnal oxygen desaturation in patients with COPD. Data exist for the cholinergic bronchodilators, ipratropium and tiotropium. Martin et al studied the effect of ipratropium inhaled four times a day in 36 patients with moderate to severe COPD (FEV\textsubscript{1} ≤ 65% predicted).\textsuperscript{82} After 4 weeks, nocturnal oxygen saturation improved, subjective sleep quality was better, and there was an increase in total REM sleep time. Tiotropium also improved nocturnal oxygen saturation, although sleep quality was not affected.\textsuperscript{83} A recent study with aclidinium bromide, a novel long-acting antimuscarinic drug, investigated its potential effects on night-time symptoms in COPD. This study showed that the improvement in lung function obtained by administration of aclidinium bromide 400 mg twice daily translated into symptomatic benefits in COPD patients, especially during sleep and in the early morning.\textsuperscript{84} Long-acting β-agonists, such as salmeterol, show similar benefits.\textsuperscript{85} Oral steroid therapy in stable COPD improves nocturnal oxygen desaturation and increases total sleep time.\textsuperscript{86} Although there are no relevant data, we might expect a similar improvement with inhaled corticosteroids. Taken together, the data suggest that treatment of COPD with or without hypercapnia will ameliorate nocturnal oxygen desaturation, and may decrease the need for supplemental oxygen as well as noninvasive ventilation. In Table 3, we outline some suggested management options for COPD patients with the desaturator phenotype who have sleep-related hypoxemia and/or hypercapnia.

### Conclusion and future perspectives

Cluster analysis may prove useful in COPD phenotyping. Although the goal of cluster analysis is to reduce the number of observations or cases by grouping them into a smaller set of clusters, the goal of factor analysis is to reduce the number of variables by grouping them into a smaller set of factors. However, these types of analyses would still require longitudinal validation to determine how such clustered subjects differ in terms of important clinical outcomes. In addition, such analyses may or may not ultimately be useful in defining specific biologic pathways or therapies.

From a practical standpoint, validation of phenotypes in COPD will require longitudinal data collection in carefully characterized patient populations. Studies such as ECLIPSE (Evaluation of COPD Longitudinally to Identify Predictive Surrogate Endpoints), that are systematically gathering clinical, physiological, radiological, biological, and genetic data on COPD subjects, will aid in this regard.\textsuperscript{87}

Our study, albeit in only 51 patients, describes a statistical methodology allowing the identification of clinical COPD desaturator phenotypes. This study shows that desaturator
patients can be identified not by their $T_{aO_2}$ value alone but by a pattern of $T_{aO_2}$, mean pulmonary artery pressure, and PaCO$_2$ values, and that the latter two variables are predictors of the severity of nocturnal desaturation. In any event, a $T_{aO_2}$ cutoff value does not appear to describe desaturator patients adequately or to assess the severity of nocturnal desaturation correctly. Moreover, cluster analysis identified subgroups of desaturator and nondesaturator patients who differ in their degree of disease severity.

We propose that dissemination of this original approach could result in better phenotypic characterization, which may prove useful in both clinical practice and in clinical trials. We further propose that data from large clinical trials should be reanalyzed using this methodology for classification of patients according to their clinical characteristics at study entry. These preliminary data would provide a clear rationale for further clinical studies. The priority would then be to determine whether the phenotypes vary in their response to different pharmacological treatments. This knowledge could lead to treatment specifically targeted at defined phenotypic groups, rather than asthma or COPD in general, which is the current management approach. Ultimately, whether different treatment strategies would provide different outcomes for these groups will confirm or refute the clinical value of cluster analysis. This knowledge could lead to different pharmacological treatments and other interventions directed at specific phenotypic groups. We consider that achieving this goal is worthy of the research endeavor.

Acknowledgments
The authors would like to acknowledge Lubello Roberto, Legari Giulia, D’Andrea Agostino, and Fausto Meleleo, for their excellent technical assistance. The authors also thank Franca Carbonaro, translator, for editing this article.

Disclosure
The authors declare that they do not have any conflicts of interest or financial relationship with any commercial entity mentioned in this work.

References


