Pathophysiological mechanism of long-term noninvasive ventilation in stable hypercapnic patients with COPD using functional respiratory imaging

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Introduction: Patients with severe COPD often develop chronic hypercapnic respiratory failure. Their prognosis worsens and they are more likely to develop exacerbations. This has major influence on the health-related quality of life. Currently, there is no information about the success of long-term noninvasive ventilation (NIV) among patients who receive NIV in acute settings. Also, little is known about the pathophysiological mechanism of NIV.

Methods: Ten Global Initiative for Obstructive Lung Disease stage III and IV COPD patients with respiratory failure who were hospitalized following acute exacerbation were treated with NIV using a Synchrony BiPAP device for 6 months. Arterial blood gases and lung function parameters were measured. Low-dose computed tomography of the thorax was performed and used for segmentation. Further analyses provided lobe volume, airway volume, and airway resistance, giving an overall functional description of the separate airways and lobes. Ventilation perfusion (VQ) was calculated. Patient-reported outcomes were evaluated.

Results: PaCO₂ significantly improved from 50.03 mmHg at baseline to 44.75 mmHg after 1 month and 43.37 mmHg after 6 months (P<0.006). Subjects showed improvement in the 6-minute walk tests (6MWTs) by an average of 51 m (from 332 m at baseline to 359 m at 1 month and 383 m at 6 months). Patients demonstrated improvement in self-reported anxiety (P=0.018). The improvement in image-based VQ was positively associated with the 6MWT and the anxiety domain of the Severe Respiratory Insufficiency Questionnaire.

Conclusion: Though previous studies of long-term NIV have shown conflicting results, this study demonstrates that patients can benefit from long-term NIV treatment, resulting in improved VQ, gas exchange, and exercise tolerance.

Keywords: noninvasive ventilation, COPD, functional respiratory imaging, ventilation perfusion, BiPAP, hypercapnia, exacerbations

Introduction

Chronic obstructive pulmonary disease (COPD) is a progressive disease that affects an increasing number of patients due to aging of the population.¹ Some patients become especially handicapped with pronounced respiratory failure, sometimes with both severe hypoxia and hypercapnia. Pharmacological treatments in this population remain controversial. Systemic steroids have been shown to be beneficial by enhancing recovery during acute exacerbations and reducing hospital stays.² ³ They are, however, not suited for long-term treatment because of severe side effects.⁴ Inhaled steroids do not slow FEV₁ decline but may prevent exacerbations in some patients, although the latter has
recently been questioned again. For these reasons, clinicians often have to rely on non-pharmacological treatments.

Among these treatments, noninvasive ventilation (NIV) stands out as a possible effective therapy. Convincing evidence has been provided over the years for its use during acute exacerbations. Prediction factors, like the severity of hypercapnia and acidosis, have been defined. Chronic use of NIV in stable hypercapnic COPD patients, however, remains very controversial. Initial studies and reviews have shown overall negative results, likely due to a lack of standardization of patients, different ventilator settings, and not well-standardized concomitant treatments. However, with increasing experience and technological advances, it has become clear that some patients may benefit from this treatment. Moreover, NIV seems to be widely used even without clear evidence for its effectiveness. During the last decade, some groups have indicated that the lack of response may be due to the use of low inspiratory positive airway pressure (IPAP). Higher pressures are shown to be more effective but limit the use because of intolerance. Given the high unmet need in severe COPD patients and the clear indication that patients can benefit, we are urged to make significant progress in this field. A different approach to solve the outstanding question of defining responders is needed. For this, a better understanding of the pathophysiological effects induced by NIV, and its influence on patient-reported outcomes (PROs), is necessary. Among the relevant pathophysiologic parameters are regional changes in airway caliber, perfusion, and ventilation perfusion ratio. The latter determines gas exchange, mainly hypoxemia, and has the potential to correlate with clinical markers such as exercise tolerance and quality of life. In a previous study, we showed that NIV can significantly redirect internal airflow presumably by opening partially or completely occluded airways, especially distal airways. When airflow is redirected toward well-perfused areas, the ventilation perfusion ratio may improve. In this study, arterial PO2 improved in some patients but regional changes in ventilation perfusion (VQ) were not measured directly.

The aim of the present study is to measure the ventilation and perfusion at the lobar level directly using functional respiratory imaging (FRI). The latter method directly measures airway caliber, airway resistance, regional flow, regional perfusion, and regional VQ. This study examined the most advanced FRI parameters to see whether these parameters changed during NIV and, more importantly, whether they correlated with PROs and exercise tolerance. This information sets the stage for algorithms that predict changes in these physiological parameters and can help identify responsive patients.

**Patients and methods**

Ten patients with Global Initiative for Obstructive Lung Disease (GOLD) stages III and IV COPD (FEV 1 <50%, Tiffeneau <70%) participated in this study. Patients were included if they were 18–80 years of age, had a diagnosis of COPD GOLD III or IV (FEV 1 <50%, Tiffeneau <70%), were hospitalized due to an exacerbation, and had developed persistent hypercapnia (PaCO2 >45 mmHg) on day 5–12 under maximal pharmacological treatment. They must have stopped smoking and must not have had any treatment with home NIV before admission. Patients who were invasively ventilated and/or had been diagnosed with asthma, restrictive lung diseases, malignancy, heart failure, or obstructive sleep apnea syndrome were excluded. The approval of the local ethical committee was obtained.

Primary outcome variables were arterial blood gas values and functional imaging of the lungs. Secondary outcome variables were lung function tests (static and dynamic lung volumes, diffusion) and exercise tolerance.

NIV was provided with a Synchrony BiPAP device (Respironics, Inc, Murrysville, PA, USA), for ≥5 hours a day, with full face mask, starting 5–12 days after admission. Modes were adapted until O2 saturation was >90% during 90% of the time and PaCO2 was decreased 5% in 1 hour. Patients were ventilated for at least 6 months and were followed up for 12 months.

Arterial blood gases were observed before starting NIV and again after 1 and 6 months. Between day 5 and 12, lung function tests and 6-minute walk tests (6MWTs) were performed and then repeated after 1 and 6 months. The following questionnaires were completed: the Saint George’s Respiratory Questionnaire, Severe Respiratory Insufficiency (SRI) Questionnaire, visual analog scale Measure of Clinical Dyspnea, and UZA BiPAP.

Low-dose high-resolution computed tomography (HRCT) scans were performed at baseline and after 1 and 6 months. One scan was taken at total lung capacity (TLC), the lung level attained after deep inspiration, and another was taken at functional residual capacity (FRC), the lung volume attained after a normal expiration. A handheld pneumotach was used for volume gating purposes to ensure scans were taken at the correct lung level.

The HRCT scans were then processed using the FRI approach which quantifies lobar volumes (iVlobes), airway volumes (iVaw), airway resistance (iRaw), and blood
vessel volumes (iVbv). FRI provides more information on regional lung function characteristics (FLUIDDA, Kontich, Belgium).14-16 HRCT images were imported into Mimics, a commercial, United States Food and Drug Administration (FDA)-approved, medical imaging processing software package (Materialise, Leuven, Belgium; FDA, K073468; CE certificate, BE 05/1191 CE01). This software package converts the HRCT images into patient-specific, three-dimensional (3D) computer models of the lung lobes and the airway tree, which can be segmented down to the bronchi with a diameter of ~1–2 mm. The 3D models were converted in a computational grid using a commercial software package (Mimics 15.0; Materialise NV, Leuven, Belgium; Food and Drug Administration, K073468; Conformité Européenne certificate, BE 05/1191.CE.01). Perfusion and ventilation were calculated separately for each lobe. Blood vessel density can be considered a surrogate for perfusion. Image-based perfusion (iQ) was calculated by blood vessel density at TLC multiplied by image volume at TLC. Image-based ventilation (iV) was calculated by the imaged volume at TLC subtracted from the image-based volume at FRC. This was calculated for the five lobes separately. The mean value of image-based ventilation perfusion match (iVQ) for each individual patient was calculated.

The study protocol was approved by the local ethics committee (Antwerp University Hospital) and written informed consent was obtained from each patient at the time of entry to the study. The study registration number is NCT01592656.

Statistical analysis
All statistical analyses were performed in R version 3.0.2 (R Foundation for Statistical Computing, Vienna, Austria). The statistical significance threshold was set to P<0.05 for all analyses.

Differences between the baseline, 1 month, and 6 month visits were assessed using linear mixed models, setting the visit as the fixed effect and the patient as the random effect. A posteriori t-test was performed to obtain the P-values between each of the two visits separately. These results are visualized by means of boxplots. In these figures, the extremes of the box represent the quartiles and the black line gives the median. The whiskers extend to the most extreme data point which is no more than 1.5 times the interquartile range from the box. All data points outside this range (outliers) are visualized as individual points. Asterisks indicate significant changes from baseline.

Correlations between clinical and FRI parameters were also assessed by linear mixed models. These results are visualized by standard x–y plots where each dot represents a single measurement. The fitted line represents the fixed effect of the linear mixed model.

Results
Patients were treated with NIV using a maximal inspiratory pressure (IPAP) of 16.3±2.4 cm H2O, a minimal inspiratory pressure of 12.0±2.4 cm H2O (average volume assured pressure support system used), and a mean expiratory pressure of 5.0±0.9 cm H2O (Table 1). Patients showed significant improvement in hypercapnia, with PaCO2 ranging from a mean of 50.03 mmHg at baseline to 44.75 mmHg after 1 month and 43.37 mmHg after 6 months (Figure 1; P=0.006). Most patients who improved after 1 month also showed improvement after 6 months. Changes in PaCO2 were not accompanied by significant changes in lung volumes. Both FVC and TLC remained almost unchanged (Figure 2; P=0.108). Although improvement in the 6MWT was not statistically significant, the mean increase of 50 m (from 332 m at baseline to 359 m at 1 month and 383 m at 6 months) is clinically meaningful (Figure 3). Statistically significant changes were also found in the anxiety domain of the SRI questionnaire (Figure 4; P=0.018).

While the changes in airway resistance and airway volume were not statistically significant, results demonstrate a trend toward an increase in airway volume and decrease in airway resistance at 6 months (Figure 5). There was a significant negative correlation between the drop in PaCO2 and inspiratory capacity (Figure 6; P=0.01). A similar correlation was seen between oxygen saturation at the end of the 6MWT and inspiratory capacity, indicating that hyperinflation goes along with more desaturation during exercise (Figure 7; P=0.03). The SRI anxiety domain score was

<p>| Table 1 IPAP and EPAP pressures used during the study |</p>
<table>
<thead>
<tr>
<th>Patient ID</th>
<th>Vt (mL)</th>
<th>bf (BPM)</th>
<th>IPAP max (cm H2O)</th>
<th>IPAP min (cm H2O)</th>
<th>EPAP (cm H2O)</th>
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<tr>
<td>1</td>
<td>350</td>
<td>12</td>
<td>16</td>
<td>12</td>
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<td>12.1</td>
<td>16.3</td>
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Abbreviations: IPAP, inspiratory positive airway pressure; EPAP, expiratory positive airway pressure; Vt, tidal volume; bf, breathing frequency.
positively associated with an improvement in distal airway volume (Figure 8; \( P < 0.001 \)). Improvement in VQ was also positively correlated with the 6MWT and the SRI anxiety domain score (Figure 9; \( P < 0.001; P = 0.04 \)).

**Discussion**

In this study, we demonstrated that NIV provided to COPD patients following a hypercapnic exacerbation lowers their PaCO\(_2\) months later without requiring adjustments to pharmacological treatments. Exercise tolerance also appears to improve with NIV therapy, especially as associated with ventilation perfusion (VQ). A drop in hyperinflation is associated with improvement in blood gases. Therefore, improvement in VQ is probably due to increased ventilation.

It has been previously hypothesized that NIV can improve VQ relationships through recruitment of poorly ventilated lung units.\(^{18}\) However, this has not been confirmed in follow-up clinical studies with relevant intermediate outcome parameters.\(^{19}\) FRI technology has made it possible to measure VQ at the lobar level in patients with hypercapnic COPD.
using long-term NIV, demonstrating tangible changes in VQ and its associated impact on relevant clinical outcomes like exercise tolerance and quality of life.

In an earlier controlled pilot study, we randomized 15 patients to a pharmacological treatment or to standard of care and NIV for 6 months.\(^1\) We assessed arterial blood gases and lung function parameters, and performed a low-dose computed tomography of the thorax with segmentation. Regional airway resistance and internal flow distribution were calculated using FRI. Blood gases, both hypoxia and hypercapnia, improved. There was a significant improvement in exercise tolerance as measured by the 6MWT. FRI showed remodeling after 6 months of treatment, with a redistribution of airflow toward well-perfused lobes in some patients, presumably due to the opening of partially or completely occluded airways, especially distally.\(^6\)

It is hypothesized that the NIV ventilator settings may be important as higher inspiratory pressures will have more capacity to open up distal airways. Specifically, the mean inspiratory pressure and backup frequencies must be high
Figure 5 A trend toward an increase in iVaw and drop in iRaw was seen when measured at TLC.

Notes: The extremes of the box represent the quartiles and the black line gives the median. The whiskers extend to the most extreme data point which is no more than 1.5 times the interquartile range from the box. All data points outside this range (outliers) are visualized as individual points. There was no significant change.

Abbreviations: TLC, total lung capacity; NIV, noninvasive ventilation; iVaw, imaged airway volume; iRaw, imaged airway resistance; m, month.

Figure 6 A significant correlation was found between a decrease in arterial PCO\textsubscript{2} (ABGpCO\textsubscript{2}) and increase in IC.

Abbreviations: ABG, arterial blood gas; IC, inspiratory capacity; TLC, total lung capacity; REML, restricted maximum likelihood.
enough to improve the alveolar ventilation and thereby reduce chronic hypercapnia.\textsuperscript{14,20} Köhnlein et al\textsuperscript{21} came to the same conclusion. They performed a large randomized controlled trial of 150 patients treated with NIV in an acute setting. They were randomized to usual standard of care or to continuing NIV for 1 year. They showed that, with effective ventilator strategies, it is possible to reduce hypercapnia significantly. This was associated with significant improvements in overall mortality (12\% vs 33\% for the controls), quality of life, and exercise capacity. In a recent study of 42 COPD patients with acute exacerbation, FRI technology demonstrated that an increase in distal airway resistance mainly led to hyperinflation and V/Q mismatch during the acute phase of an exacerbation.\textsuperscript{22} In the recovery phase, there was significant improvement in distal resistance and improvement in VQ.

Collectively, these studies suggest that exacerbation treatment should therefore be focused on improvement of distal airway resistance. By increasing the intraluminal airway pressure with NIV, distal airway resistance will reduce, providing a better VQ match, especially due to an increase in the ventilation related to a drop in distal airway resistance. Opening the distal airways with systemic therapy,\textsuperscript{23} inhaled therapy, and/or airway clearance techniques\textsuperscript{24–29} are adequate
treatments for COPD exacerbation and persistent respiratory failure due to COPD.

Lastly, improvement in distal airway resistance can also lead to improved PROs and can therefore be considered as a useful surrogate clinical marker. In this study, patients improved their exercise capacity with an average increase of 51 m in the 6MWT. Self-reported anxiety also improved over the course of treatment. These findings demonstrate that the observed changes in physiological parameters have a clinical meaning.

There has been a long debate about the usefulness of NIV in the chronic treatment of COPD. While some overviews argue against the chronic use of NIV in these patients, this study demonstrates that patients can benefit from treatment via improved VQ leading to improved gas exchange and thereby better exercise tolerance. However, not all patients respond to this treatment. Nevertheless, this is not a reason to withhold treatment from all patients, especially when we now are realizing that responders can have significant benefit.

FRI allows us to measure relevant surrogate physiological markers, like VQ, and correlate them with clinical outcomes. It offers the possibility of detecting responders in early phases of treatment and establishing stopping rules.

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Disclosure
J De Backer, W Vos, F Ferreira, and KC Barboza are affiliated with Fluidda NV. The authors have no other relevant
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References