Noninvasive Oxygenation in Patients with Acute Respiratory Failure: Current Perspectives

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Purpose of Review: High-flow nasal oxygen and noninvasive ventilation are two alternative strategies to standard oxygen in the management of acute respiratory failure.

Discussion: Although high-flow nasal oxygen has gained major popularity in ICUs due to its simplicity of application, good comfort for patients, efficiency in improving oxygenation and promising results in patients with acute hypoxemic respiratory failure, further large clinical trials are needed to confirm its superiority over standard oxygen. Non-invasive ventilation may have deleterious effects, especially in patients exerting strong inspiratory efforts, and no current recommendations support its use in this setting. Protective non-invasive ventilation using higher levels of positive-end expiratory pressure, more prolonged sessions and other interfaces such as the helmet may have beneficial physiological effects leading to it being proposed as alternative to high-flow nasal oxygen in acute hypoxemic respiratory failure. By contrast, non-invasive ventilation is the first-line strategy of oxygenation in patients with acute exacerbation of chronic lung disease, while high-flow nasal oxygen could be an alternative to non-invasive ventilation after partial reversal of respiratory acidosis. Questions remain about the target populations and non-invasive oxygen strategy representing the best alternative to standard oxygen in acute hypoxemic respiratory failure. As concerns acute on-chronic-respiratory failure, the place of high-flow nasal oxygen remains to be evaluated.

Keywords: noninvasive ventilation, high-flow nasal oxygen, respiratory insufficiency, acute respiratory failure

Introduction

Respiratory failure is defined as the failure of the respiratory system in one or both of its gas exchange functions, ie, oxygenation of and/or elimination of carbon dioxide from mixed venous blood.1 Oxygen therapy is usually the first-line treatment in acute respiratory failure and classically includes standard oxygen, noninvasive ventilation (NIV) and, more recently, high flow nasal cannula oxygen therapy (HFNC). Oxygen therapy delivered through a non-rebreathing face mask is the oldest non-invasive oxygen support, first described in 1946.2 NIV was developed less than 10 years after, in 1952, in an initial study reporting NIV delivered through continuous positive airway pressure (CPAP) in cardiogenic pulmonary edema.3 After the 1990s, NIV through CPAP or applied with pressure support (PS) and positive end expiratory pressure (PEEP) have been widely used with a strong level of evidence in cardiogenic pulmonary edema4 and COPD exacerbation.5,6 HFNC is another noninvasive oxygen support, first described in 1968, and is known for its ability to provide a high level of FiO2 due to high gas flow.7 In the 2010s, its use spread among adult patients with acute hypoxemic respiratory failure, after first having been used in pre-term neonates and pediatric care, as a first-line treatment for respiratory distress syndrome, and apnea of prematurity. Contrary to NIV, most of the clinical studies on HFNC have preceded physiological studies and reported good comfort and a better prognosis in acute hypoxemic respiratory failure than with other noninvasive oxygenation supports, thereby justifying its widespread use in intensive care units.8,9

The expected aims of noninvasive oxygenation supports (Table 1) are to provide comfort by preserving physiological pathways of airway protection (ie, cough and clearance of secretions)10 and to avoid invasive mechanical ventilation and related morbidity or mortality. Adverse effects include ventilator-induced lung injury,11,12 ventilator-associated...
pneumonia, and complications related to sedation and neuromuscular blockage. Spontaneous breathing may have further benefits such as preventing diaphragm dysfunction and atrophy, and it yields increased aeration of the dependent lung, which minimizes ventilation/perfusion mismatch. However, maintenance of spontaneous breathing may expose the patient to prolonged inspiratory efforts and delayed intubation with potential impact on prognosis. This review summarizes existing literature about the efficacy and safety of non-invasive oxygenation supports, ie, NIV and HFNC, in the management of acute respiratory failure.

**Acute Hypoxemic Respiratory Failure**

Acute hypoxemic respiratory failure or de novo respiratory failure is characterized by a gas exchange failure manifested by severe acute hypoxemia without hypercapnia. Hypoxemia results from lung failure due to an unequal ventilation/perfusion ratio, and increased shunt and/or diffusion impairment. The definition of acute hypoxemic respiratory failure is not clearly established. The level of hypoxemia varies from study to study with a PaO$_2$/FiO$_2$ ratio equivalent below or equal to 300 mm Hg or 200 mm Hg. The most common cause is pneumonia, and acute-on-chronic respiratory failure and cardiogenic pulmonary edema are ruled out. Acute respiratory distress syndrome (ARDS) is a subset of acute hypoxemic respiratory failure. However, to fulfill the definition, it requires not only the presence of bilateral lung infiltrates on chest imaging but also a positive end-expiratory pressure (PEEP) level of at least 5 cmH$_2$O assessed under positive pressure ventilation and hypoxemia not fully explained by fluid overload or cardiac dysfunction.

During acute hypoxemic respiratory failure, breathing patterns result in hyperventilation due to high respiratory drive and inspiratory effort leading to high tidal volumes, increased respiratory rates and, finally, to hypocapnia. However, spontaneous breathing in this setting can be potentially harmful and lead to a vicious circle generated by hypoxemia, dysregulated and high inspiratory effort, leading in turn to transpulmonary pressure swings and potential inhomogeneous redistribution of tidal volume generating local injurious forces. The concept of patient self-inflicted lung injury (P-SILI) helps to explain ventilation-induced lung injury during spontaneous breathing (Figure 1).

**Physiological Responses of Non-Invasive Oxygen Support**

Consequently, one of the objectives of non-invasive oxygen support is to unload respiratory effort and to mitigate the consequence of high inspiratory effort (Table 1). However, such a strategy should be conducted cautiously so as to avoid delayed intubation, which may increase risk of mortality. Since the end of the 90's, NIV has shown contradictory results in acute hypoxemic respiratory failure, while HFNC is currently on the rise after having delivered benefits in patients with acute hypoxemic respiratory failure. HFNC is an oxygen device able to deliver high flows of warmed humidified gas, up to 70 L/min with a temperature set from 33° to 37° C, with a FiO$_2$ ranging from 21% to 100%. Physiological responses to HFNC include PEEP effect, due to low levels of positive pressure in the upper airways favoring an increase in end-expiratory lung volume and oxygenation, and also to ventilatory support due to the effects on dead-space washout, leading to a decreased respiratory rate and, finally, work of breathing (Figure 1). During acute hypoxemic respiratory failure, the inspiratory effort leads to a high peak inspiratory flow reaching a mean of 30–40 L/min, which can exceed 60 and even reach 120 L/min in more severe patients. The high flow generated by the HFNC system is able to provide high and controlled FiO$_2$. Although the system is open, delivery of a high flow rate usually set at 50 L/min and able to reach 70 L/min in most cases exceeds the patient’s peak inspiratory flow rate and preserves from dilution of inhaled gas with room air. In a physiological study including healthy

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**Table 1 Aims of Noninvasive Oxygen Supports in Acute Respiratory Failure**

<table>
<thead>
<tr>
<th>Acute hypoxemic respiratory failure</th>
<th>To avoid intubation and to decrease related mortality</th>
<th>Acute hypercapnic respiratory failure</th>
</tr>
</thead>
<tbody>
<tr>
<td>To preserve tolerance</td>
<td>Comfort</td>
<td>To preserve tolerance</td>
</tr>
<tr>
<td>PaO$_2$, PaO$_2$/FiO$_2$</td>
<td>Blood gas improvement</td>
<td>PaCO$_2$</td>
</tr>
<tr>
<td>Inspiratory effort</td>
<td>To unload inspiratory muscles</td>
<td>Inspiratory effort</td>
</tr>
<tr>
<td>To ensure prevention of lung injury</td>
<td></td>
<td>Expiratory flow limitation</td>
</tr>
</tbody>
</table>

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volunteers, Sim et al measured FiO\textsubscript{2} in their pharynx during oxygen delivery through several devices such as a standard mask, a non-rebreathing mask and HFNC system\textsuperscript{23}. With a standard mask, FiO\textsubscript{2} did not exceed 0.6 despite a flow of 12 L/min, and it dropped below 0.5 when acute respiratory failure was simulated by thoracic contention. Although the non-rebreathing mask avoided such a FiO\textsubscript{2} drop, the highest FiO\textsubscript{2} obtained under an oxygen flow rate of 15 L/min was less than 0.7. By comparison, FiO\textsubscript{2} reached 0.85 using the HFNC system set, with a flow rate of 40 L/min\textsuperscript{23}. Another effect of the HFNC system is to generate a low level of positive pressure in the upper airway, which is directly proportional to the gas flow delivered. However, due to the open nature of the system (with air leakage), the pressure levels are quite variable and not controlled. Although positive pressure is markedly reduced when the patient opens his mouth, it generates a PEEP effect with alveolar recruitment that may help to improve gas exchange. In physiological studies assessing pulmonary volumes with electrical impedance tomography after cardiac surgery or during acute hypoxemic respiratory failure, increased end-expiratory lung volume directly proportional to gas flow\textsuperscript{24} was found with HFNC as compared to standard oxygen therapy, suggesting alveolar recruitment reflecting a PEEP effect.\textsuperscript{25,26} Given these different physiological effects, HFNC can appear as a support for oxygenation able to decrease work of breathing. Many authors have highlighted in patients with acute respiratory failure treated first by standard oxygen through a non-rebreathing mask and then by HFNC, a decrease in work of breathing under HFNC assessed by swings of esophageal pressure (diminution of pressure–time product during inspiration).\textsuperscript{26,27} Mauri et al assumed that better working conditions could be partially due to improvement of inspiratory effort and pulmonary compliance possibly favored by increased ventilation homogeneity.\textsuperscript{26} Otherwise, the high-flow rate of continuously delivered gas may flush the upper airways, generating a washout of dead space and then flushing out carbon dioxide.\textsuperscript{28,29} This effect, which is associated with mechanically improved thoracic properties, results in reduced inspiratory effort and minute ventilation requirement. This is consistent with the commonly reported finding of decreased respiratory rate and work of breathing with HFNC.\textsuperscript{26,27} Consequently, HFNC improves pattern of breathing by increasing oxygenation,
decreasing inspiratory effort, respiratory rate and work of breathing, thereby possibly mitigating the consequences of self-inflicted lung injury (Figure 1).

The main characteristic of NIV is assistance of spontaneous breathing by administration of preset positive pressure into the lungs through an external interface, ie, a commonly used face mask connected to a humidification system, a heated humidifier or a heat and moisture exchanger, and a ventilator. The most commonly used mode combines pressure support (PS) ventilation with PEEP, or else simply applies continuous positive airway pressure (CPAP). NIV through facemask is usually applied with low PEEP levels (5–8 cmH2O), because of the presence of air leaks potentially leading to discomfort and poor tolerance. NIV is able to increase oxygenation by providing high FiO2 up to 100% and through PEEP and to achieve gas exchange as effective as invasive ventilation. In addition to improving oxygenation, PEEP alone or CPAP reduces inspiratory effort in spontaneous breathing patients with acute respiratory failure compared to standard oxygen, as attested by a reduction in esophageal pressure swings. Moreover, Morais et al showed that increasing PEEP levels minimized lung injury, especially in dependent lung, in animals with ARDS and spontaneous breathing under mechanical ventilation. In a physiological study including patients under NIV for mild ARDS, increased pressure support resulted in proportionally decreased inspiratory effort and dyspnea, as has been determined by changes in esophageal and diaphragmatic pressures. Moreover, increasing pressure support resulted in proportionally increased tidal volumes and potential alveolar recruitment, as suggested by some authors (Figure 1).

However, air leaks, discomfort and skin breakdown limit the tolerability of facemask NIV, rendering prolonged treatments with specific settings, ie high PEEP, difficult to apply, and it may even lead to a need for intubation in 10% of the cases. Several authors have proposed the helmet interface as an alternative interface for NIV delivery in hypoxemic patients. The helmet is a transparent hood that covers the entire head, sealed with a soft neck ring, thereby producing a breathing circuit semi-closed from the outside environment. NIV delivered through helmet has been shown to provide better tolerability and less air leaks, enabling the possibility to deliver prolonged treatments with high PEEP reaching 12 cmH2O. A physiological study comparing NIV delivered through helmet and face mask showed that with a helmet, improved breathing patterns, especially inspiratory muscle load, consisted in high PEEP and pressure support levels, which increased by 50% compared with the facemask settings. Indeed, some of the pressure support is dissipated in the helmet and does not necessarily correspond to the pressure inside at the airway opening and in the alveoli. Therefore, the optimization of NIV delivery may involve changes in ventilator settings by increasing PEEP levels and changes in the interface improving comfort, with the aim of achieving prolonged sessions.

**Clinical Impact and Limitations of Non-Invasive Oxygen Supports**

As reported with invasive mechanical ventilation, conventional NIV delivered with facemask may be deleterious in acute hypoxemic respiratory failure due to barotrauma favored by the high respiratory drive, leading to high inspiratory effort, and the synchronization with the pressure-support, which may result in high tidal volumes. Accordingly, Tonelli et al have put forward the hypothesis of a relationship between excessive spontaneous patient effort and NIV failure. In the 30 included patients with acute hypoxemic respiratory failure undergoing a trial of NIV, tidal changes in esophageal pressure were not reduced within 2 hours in patients who failed with NIV, whereas these changes were significantly reduced in those who succeed with NIV. High expired tidal volumes, ie, exceeding 9.5 mL/kg of predicted body weight, were independently associated with NIV treatment failure. Similarly, two clinical studies had previously reported that tidal volumes exceeding 9 or 9.5 mL/kg of predicted body weight during NIV application were strongly associated with intubation and mortality in larger samples of patients treated for acute hypoxemic respiratory failure.

In the study by Carteaux et al, the authors attempted to target a tidal volume from 6 to 8 mL/kg of predicted body weight under NIV, while nearly half of the patients had a tidal volume above 10 mL/kg. In our study, NIV settings between intubated and not intubated patients were similar, with pressure support around 8 and PEEP around 5 cmH2O, notwithstanding the higher tidal volumes generated in intubated patients. Patients who needed intubation under NIV did not exhibit higher severity scores or higher respiratory rates than the others, ie the usual criteria to assess severity at bedside. Bellani et al showed that during spontaneous breathing in patients receiving invasive ventilation, a significant reduction of pressure support did not result in decreased patient-generated tidal volume and did not reduce transalveolar pressure. Therefore, high tidal volumes might be considered as the consequence of the high respiratory drive leading to...
high inspiratory efforts due to hypoxemia, and thereby reflect the severity of the underlying respiratory disease. Moreover, they may further worsen a pre-existing lung insult by inducing ventilator-induced lung injury (VILI). The clinical benefits of conventional NIV delivered through facemask in patients with acute hypoxemic respiratory failure are uncertain\(^8\)\(^{,47–51}\) as underlined in recent clinical practice guidelines on the use of NIV.\(^6\) Indeed, international experts were unable to recommend NIV in this setting, given the low certainty of evidence concerning risks of intubation and mortality.\(^6\)

However, NIV delivered through helmet could change this paradigm with different settings and subsequent physiological effects. In a monocenter randomized controlled trial including 83 patients with ARDS treated first with NIV through facemask, NIV delivered secondarily through helmet seemed to be more beneficial than NIV continued through facemask, ie, lower intubation and mortality rates.\(^52\) In addition to different interface application, NIV settings significantly differed between groups with higher positive end-expiratory pressure and lower pressure support levels in the helmet group. Benefits were probably due to the physiological advantages of helmet, namely delivery of higher PEEP in continuous sessions with enhanced comfort. Whether the benefits are due to decreased inspiratory effort or to optimized alveolar recruitment is not clear. In fact, measurement of tidal volumes under helmet, ie, a simple approach of inspiratory effort, is not routinely feasible and did not appear in the trial. As a result, the impact of settings and potential contribution to reduced transpulmonary pressure and lung injury (VILI) by decreasing ventilator assistance is difficult to analyze.\(^52\)

Whether HFNC is efficient in preventing superimposed lung injury in spontaneous breathing patients during acute hypoxemic respiratory failure remains to be established. In a randomized controlled trial comparing NIV, HFNC and standard oxygen in 313 patients with acute hypoxemic respiratory failure, we reported significant differences in favor of HFNC in terms of mortality (12% versus 23% and 28% with standard oxygen and NIV, respectively) and of intubation for severe hypoxemic patients (38%, 47% and 50%, respectively).\(^8\) Considering these results, HFNC appeared superior to NIV and standard oxygen, given that it could prevent VILI (as compared to NIV) and also PSILI (as compared to standard oxygen). However, in another large randomized controlled trial including 778 immunocompromised patients with acute respiratory failure, the superiority of HFNC to standard oxygen was not confirmed as regards risk of intubation (39% versus 44%, respectively) or mortality (36% in both groups of treatment).\(^53\) Accordingly, recent clinical practice guidelines\(^54\) and a meta-analysis\(^55\) reported a reduced risk of intubation but not of mortality with HFNC as compared to standard oxygen. Currently, there is no strong evidence of the superiority of HFNC or conventional NIV delivered through facemask over standard oxygen in the management of acute hypoxemic respiratory failure whatever the severity. Although HFNC has gained popularity among intensivists thanks to its simplicity of application, good comfort for patients and efficiency in improving oxygenation, only one randomized controlled trial has reported benefits of HFNC as compared to NIV or standard oxygen,\(^8\) whereas another randomized controlled trial did not show any superiority of HFNC as compared to standard oxygen.\(^53\) Further studies are needed to confirm the place of HFNC in this setting and could help to determine the patient population, ie, mild, moderate or severe hypoxemic patients, most likely to benefit from HFNC in terms of intubation and mortality.

Experience of the Pandemic: Optimization of Non-Invasive Oxygen Support Strategies

During the COVID-19 pandemic, use of non-invasive oxygen supports was heterogeneous across countries; for instance, HFNC was less frequent in Italy or North America than in France (with use reaching 19%), while standard oxygen was the most frequent oxygen support applied.\(^56–58\) This was due to contradictory recommendations and a paucity of evidence-based management guidelines.\(^59\) One of the major concerns was to limit the spread of infection among healthcare workers, given that procedures liable to disperse viral particles, the so-called “aerosol-generating procedures”, were avoided whenever possible. Thereafter, several simulation studies using manikin model of exhaled air dispersion distances and analyzing concentrations of aerosol from the respiratory tract in room air showed that this risk was not higher under HFNC than NIV or standard oxygen devices.\(^60,61\) Moreover, barrier procedures and, specifically, the use of a surgical mask on HNFC helped to reduce the risk of bio-aerosol dispersion without impairment of patient blood oxygenation.\(^62\)
Whether or not the specificity of patients with COVID-19-induced respiratory failure might modify the efficiency of the different non-invasive oxygen supports has yet to be conclusively determined. Of note, these patients present with a remarkable disconnect at rest between profound hypoxemia without proportional signs of respiratory distress, no sensation of dyspnea or increased respiratory work, whatever the rapidity of deterioration. This is illustrated by the comparing patient populations with acute hypoxic respiratory failure caused mainly by either bacterial pneumonia and with COVID-19. Despite similar intubation rates, 38% and 34%, in the two respective populations, PaO₂/FiO₂ ratio at enrollment in non-COVID patients was higher (150–160 mm Hg) with a high respiratory rate (around 33 breaths/min), while patients with COVID-19 had more severe oxygen impairment (PaO₂/FiO₂ ratio of 102–105 mm Hg) with a surprisingly lower respiratory rate (28 breaths/min). This peculiar pattern of COVID-19-induced respiratory failure references the concept of “silent” or “happy” hypoxemia. Possible pathophysiological mechanisms include 1) intrapulmonary shunting, due to local interstitial edema, resulting in ventilation-perfusion ratio mismatch and in an alveolar to arterial oxygen gradient; 2) loss of lung perfusion regulation, with involvement of the renin-angiotensin system, intravascular microthrombi, favored by local acute inflammation; and 3) endothelial injury resulting in an imbalance between procoagulant and fibrinolytic activity; these different abnormalities lead to impaired diffusion capacity.

Silent hypoxemia during COVID-19-induced respiratory failure raises questions about the timing of initiation of non-invasive oxygen supports, whether before or after onset of tachypnea, and intubation to switch to invasive ventilation, based on level of hypoxemia or signs of respiratory distress. One potential risk of non-invasive oxygen supports is to delay intubation by masking signs of respiratory distress. A North American study of six COVID-19-specific ICUs included 231 patients, of whom 175 were intubated after having received different non-invasive oxygen supports. Timing of intubation was not associated with mortality, and patients under HFNC intubated after 24 hours of treatment did not have a higher risk of mortality than those intubated earlier. Two other observational studies have shown less intubation with HFNC as compared to standard oxygen, but no difference was reported in mortality rates. Up until now, most published studies in this setting have been observational and mainly showed better oxygenation under HFNC than standard oxygen, with intubation rates ranging from 30% to 50%. A recent trial compared HFNC with standard oxygen in 220 patients with severe COVID-19 for the need for intubation and time to clinical recovery until day 28. Authors reported that use of HFNC significantly decreased need for mechanical ventilation support and time to clinical recovery compared with standard oxygen therapy, but there was no difference in mortality between treatments, possibly due to an underpowered study. Thereby, the question of the benefits of HFNC as a treatment for the management of COVID-19 induced respiratory failure remains incompletely unanswered.

A mega-trial pooling several trials from different countries aimed to evaluate whether HNFC optimized with sessions of awake prone position in patients with severe COVID-19-related acute hypoxemic respiratory failure could reduce either death or intubation. Among the 1121 patients included in the intention-to-treat analysis, 94% were enrolled in three (Mexico, USA and France) of the six countries participating in the mega-trial. Results showed a favorable effect of awake prone position on the primary outcome, significant reduced intubation rates, while mortality rates did not differ between the two groups. However, there was heterogeneity among the different countries, with a significant effect of awake prone position in Mexico, which was not the case in the other countries. One explanation may be the longer duration of awake prone position, around 8 hours in Mexico as compared to fewer than 5 hours in the other countries. As a result, this approach requires confirmation in future studies as does the reference treatment for oxygenation.

Given the limitations of standard NIV delivered through facemask in acute hypoxemic respiratory failure, some authors have proposed protective NIV delivered through helmet, the objectives being to reduce inspiratory effort and to render spontaneous breathing less injurious by preventing the risk of VILI and P-SILI. Grieco et al showed in a physiological study that NIV delivered through helmet with a high level of PEEP (≥10 cm H₂O) and pressure support around 10 cm H₂O compared to HFNC could improve oxygenation, relieve dyspnea and reduce inspiratory effort. In 2020, the same authors conducted a multicenter randomized controlled trial, including 109 patients with severe COVID-19-related respiratory failure randomly assigned to receive HFNC at 60 L/min or NIV through a helmet set as described above. There was no difference in the primary outcome (days free of respiratory support at day 28). Although intubation rate was lower and invasive ventilation-free days at day 28 were higher after treatment by NIV through
helmet, mortality did not differ compared to HFNC. In fact, the mortality rate was higher in patients who failed helmet NIV as compared those who failed HFNC.

As a result (Table 2), HFNC may represent an alternative to standard oxygen or conventional NIV through facemask in acute hypoxemic respiratory failure. However, there is no strong evidence favoring the use of HFNC in this setting, as it seems to reduce risks of intubation but not those of mortality. While the experience of COVID-19-related respiratory failure has highlighted the potential benefit of protective NIV delivered through a helmet with high levels of PEEP compared to HFNC, it once again seems to reduce risks of intubation but not those of mortality. We are conducting a trial in France with the aim at assessing the efficiency of HFNC versus standard oxygen and at determining the standard non-invasive oxygen strategy most apt to manage patients with acute hypoxemic respiratory failure, including those infected by COVID-19 (NCT 04468126). Further studies are necessary to confirm the possible superiority of HFNC over standard oxygen and the benefits of protective NIV through a helmet.

### Acute-on-Chronic Respiratory Failure

Acute respiratory failure resulting in acute hypercapnia appears in cases of underlying chronic respiratory failure, ie chronic obstructive pulmonary disease (COPD) exacerbation. Indeed, many of these patients may have hypercapnia at baseline and the development of acidosis indicates acute-on-chronic respiratory failure. Acute respiratory acidosis occurs when the respiratory muscles fail to achieve adequate alveolar ventilation contrasting with high levels of diaphragmatic activity. Consequently, pump failure leads to increased arterial carbon dioxide (CO$_2$) and respiratory acidosis ensues.$^{1,6}$ As of now, the recommended treatment includes NIV, which provides ventilatory support, avoids intubation and a need for invasive ventilation. Since 1990, NIV has shown benefits in the management of COPD patients. An initial physiological study conducted in 1990 reported decreased PaCO$_2$ and increased pH after a 45-min NIV session.$^{70}$ Moreover, some authors have reported that diaphragmatic activity decreased significantly, when assessed either by transdiaphragmatic pressure changes or electromyographic signals.$^{70}$ Thereafter, a clinical study reported in a larger sample size of patients a reduced need for intubation and reduced hospital mortality.$^{71}$ Accordingly, the latest clinical practice guidelines including a large meta-analysis published in 2016 showed that with NIV, intubation rates have significantly decreased from around 40% with standard oxygen to 10–15% with NIV in patients treated for COPD exacerbation with hypercapnic acidosis, while mortality rates decreased from 20% to less than 10% with NIV.$^6$

As far as NIV outcomes are concerned, the current issue does not seem to further reduce the need of intubation or mortality, as these events are not common when using NIV, but rather to reduce the duration of NIV, and to offer another, more comfortable device to patients with poor tolerance to NIV or to patients with non-severe COPD exacerbation (Table 1). HFNC could be considered as ventilatory support in COPD patients, as physiological studies suggest its

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**Table 2 Summary of Proposals for First-Line Therapies in Acute Respiratory Failure**

<table>
<thead>
<tr>
<th></th>
<th>Standard Oxygen</th>
<th>Noninvasive Ventilation with Face Mask</th>
<th>Noninvasive Ventilation with Helmet</th>
<th>High-Flow Nasal Cannula Oxygen Therapy</th>
<th>Comment</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute hypoxemic respiratory failure</td>
<td>+</td>
<td>-</td>
<td>-</td>
<td>+</td>
<td>One RCT has reported benefits of HFNC versus NIV and standard oxygen. NIV is recommended by the recent European/American guidelines. However, studies are ongoing to determine the place of HFNC.</td>
</tr>
<tr>
<td>Immunocompromised patients</td>
<td>+</td>
<td>+</td>
<td>No trial</td>
<td>+</td>
<td>NIV is strongly recommended by the recent European/American guidelines. HFNC can be an alternative in case of intolerance to NIV.</td>
</tr>
<tr>
<td>COPD exacerbation with acute respiratory acidosis</td>
<td>+++</td>
<td>?</td>
<td>?</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Non-severe COPD exacerbation</td>
<td>+</td>
<td>Not recommended</td>
<td>?</td>
<td>+</td>
<td></td>
</tr>
</tbody>
</table>

**Abbreviations:** COPD, chronic obstructive pulmonary disease; HFNT, high-flow nasal cannula oxygen therapy; NIV, noninvasive ventilation; RCT, randomized controlled trial.
favorable effects on the work of breathing and gas exchange.\textsuperscript{72–74} HFNC delivers a humidified and heated gas through a nasal cannula at high flow rate, reducing anatomical dead space in the upper airways by clearing exhaled carbon dioxide.\textsuperscript{28,29} Moreover, the high flow rate generates a low level of positive pressure in the upper airways, which can provide a slight positive end-expiratory pressure effect.\textsuperscript{26,75,76} All of these physiological effects have been shown to provide help in decreasing PaCO\textsubscript{2}, respiratory rate and in improving breathing pattern with higher tidal volumes, in stable COPD patients\textsuperscript{72–74,77,78} as well as in unstable patients.\textsuperscript{81} According to Spoletini et al assessed NIV interspersed with HFNC sessions in patients treated for acute respiratory failure.\textsuperscript{81} However, this study, which included 57 patients, did not show any difference in the time spent on NIV or time on break between patients receiving standard oxygen in between NIV sessions and those receiving HFNC in between NIV sessions.\textsuperscript{81} HFNC was associated with better comfort and improved respiratory rate and dyspnea compared to standard oxygen between NIV sessions. The authors were very cautious in their conclusions insofar as their study was a non-inferiority trial with a primary physiologic outcome, leaving uncertainty on stronger patient-related outcomes. One potential limitation in explaining the cause of the failed trial is a heterogeneous population including not only patients with severe acute exacerbation of COPD but also those with acute hypoxemic respiratory failure. In this latter group, the benefit of NIV is controversial and has been reported to be associated with poor prognosis.\textsuperscript{6,8} Another ongoing trial is focused on patients requiring NIV for hypercapnic acute respiratory failure due to severe COPD exacerbation. The primary outcome is to determine whether HFNC as compared with standard oxygen increases the number of ventilator-free days (NIV and invasive mechanical ventilation) and alive at day 28.\textsuperscript{82}

One use for HFNC may be in patients with non-severe acute exacerbation of COPD. A recent trial by Cortegiani et al reported non-inferior reduction of PaCO\textsubscript{2} in patients with mild-to-moderate acute exacerbation of COPD treated with HFNC or NIV as initial ventilatory support.\textsuperscript{83} However, one-third of the patients receiving HFNC required NIV within 6 hours, and intubation rates between groups were similar.\textsuperscript{83} In less severe patients, a prospective randomized controlled trial including 320 patients with acute compensated hypercapnic respiratory failure showed a decreased intubation rate after treatment by HFNC as compared with conventional oxygen therapy, 10% versus 19%, respectively.\textsuperscript{84} Bronchodilator therapy, the main pharmacological treatment in patients having COPD exacerbation,\textsuperscript{85,86} can be delivered through HFNC circuit. Clinical studies conducted in stable ambulatory COPD patients have shown that bronchodilator therapies are effectively delivered within a HFNC circuit, providing bronchodilation similar to standard mask jet nebulization.\textsuperscript{87,88} Braunlich et al reported in 26 stable COPD patients a 9% increase in FEV\textsubscript{1} 30 minutes after bronchodilator (salbutamol and ipratropium) nebulization using a jet-nebulizer adapted on a HFNC system.\textsuperscript{88} Reminiac et al highlighted a 16% greater increase of FEV\textsubscript{1} using a vibrating nebulizer through HFNC system in 25 ambulatory patients with reversible obstructive pulmonary disease.\textsuperscript{87} In a physiological crossover study including patients with severe acute exacerbation of COPD, we showed significantly improved FEV\textsubscript{1} (9%), FVC (14%) and PEF (20%) after salbutamol vibrating-mesh nebulization through HFNC as compared to HFNC alone, a finding suggesting an effective bronchodilator effect.\textsuperscript{89}

As a result (Table 2), NIV remains the first-line therapy for severe COPD exacerbation with acidosis, whereas there is no high-level evidence favoring HFNC as first-line therapy or in less severe patients; that said, it can represent an alternative oxygen support in case of NIV intolerance with possible delivery of bronchodilator therapy.

**Conclusion**

Although HFNC is widely used in ICUs, further studies are needed to confirm its superiority over standard oxygen in acute hypoxemic or in acute-on-chronic respiratory failure, by determining the target population of patients likely to benefit from HFNC. Moreover, the benefits of NIV delivered through a helmet on HFNC in patients with acute hypoxemic respiratory failure reported in physiological studies have not been confirmed in clinical studies in terms of survival. Consequently, questions still remain about the best non-invasive oxygen strategy as an alternative to standard oxygen in acute hypoxemic respiratory failure; as concerns acute-on-chronic respiratory failure, the place of HFNC has to be evaluated.
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**References**


