Utilization of pain and sedation therapy on noninvasive mechanical ventilation in Korean intensive care units: a multi–center prospective observational study

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**Background:** The use of sedative drugs may be an important therapeutic intervention during noninvasive ventilation (NIV) in intensive care units (ICUs). The purpose of this study was to assess the current application of analgosedation in NIV and its impact on clinical outcomes in Korean ICUs.

**Methods:** Twenty Korean ICUs participated in the study, and data was collected on NIV use during the period between June 2017 and February 2018. Demographic data from all adult patients, NIV clinical parameters, and hospital mortality were included.

**Results:** A total of 155 patients treated with NIV in the ICUs were included, of whom 26 received pain and sedation therapy (sedation group) and 129 did not (control group). The primary cause of ICU admission was due to acute exacerbation of obstructed lung disease (45.7%) in the control group and pneumonia treatment (53.8%) in the sedation group. In addition, causes of NIV application included acute hypercapnic respiratory failure in the control group (62.8%) and post-extubation respiratory failure in the sedation group (57.7%). Arterial partial pressure of carbon dioxide (PaCO2) levels before and after 2 hours of NIV treatment were significantly decreased in both groups: from 61.9 ± 23.8 mm Hg to 54.9 ± 17.6 mm Hg in the control group (P < 0.001) and from 54.9 ± 15.1 mm Hg to 51.1 ± 15.1 mm Hg in the sedation group (P = 0.048). No significant differences were observed in the success rate of NIV weaning, complications, length of ICU stay, ICU survival rate, or hospital survival rate between the groups.

**Conclusions:** In NIV patients, analgosedation therapy may have no harmful effects on complications, NIV weaning success, and mortality compared to the control group. Therefore, sedation during NIV may not be unsafe and can be used in patients for pain control when indicated.

**Key Words:** intensive care units; noninvasive ventilation; safety; sedatives
INTRODUCTION

Noninvasive positive pressure ventilation is increasingly being used for the treatment of patients with acute respiratory failure [1]. The clinical benefits of this therapeutic technique have proven efficacious in various respiratory failure conditions: the acute respiratory acidosis in exacerbations of chronic obstructive lung disease (COPD); the hypoxemia derived from cardiogenic pulmonary edema [2]; the hypoxic respiratory failure observed in immunocompromised patients [3]; and in patients with advanced neuromuscular disorders such as amyotrophic lateral sclerosis. Despite the clear advantages of noninvasive ventilation (NIV) when used in critically ill patients, it remains associated with large failure rates due to patient discomfort or refusal. Recent studies have shown that NIV failure rates are approximately 40%, which remains a challenging barrier for the implementation of this effective technique [4].

Analgesia-based sedation is a common intervention used to increase the tolerance of patients who have to be supported using invasive ventilation. Similarly, there has been an effort to reduce the discomfort and refusal of patients utilizing NIV. Conti et al. [5] showed that the continuous infusion of sufentanil in patients undergoing NIV had no significant effects on respiratory variables or hemodynamic changes. However, sufentanil is a long-acting opioid, and the long-term infusion may induce the accumulation phenomena, which can delay patient recovery and augment the risk of respiratory depression [6]. Recent studies have demonstrated the safety and efficacy of analgosedation in patients with NIV failure, using drugs such as remifentanil, dexmedetomidine, and propofol [7-11]. Nonetheless, clinicians hesitate to use these analgosedative pharmacologics as they may also cause respiratory depression in NIV patients.

The aim of this study was to assess the current application of analgosedation in NIV and its impact on clinical outcomes. As such, we conducted a multicenter prospective observational study in Korean intensive care units (ICUs) which employ the NIV technique.

MATERIALS AND METHODS

Study Population

Twenty ICU intensivists belonging to Korean national hospitals participated in the study and collected data on NIV use during the period between June 2017 and February 2018. Adult patients (aged > 18 years) who were admitted to these ICUs and received NIV treatment for acute respiratory failure were prospectively included. This data was one part of a nationwide data collection study on NIV in Korean ICUs [12]. The indications for NIV included the following: (1) acute hypercapnic respiratory failure, which was designated as respiratory failure occurring in a patient with chronic lung disease (obstructive or restrictive) and, (2) de novo respiratory failure, typically indicating respiratory failure in patients without chronic respiratory disease, primarily found in those with hypoxic respiratory failure (e.g., pneumonia, postoperative respiratory failure, sepsis, or acute respiratory distress syndrome) [13,14]. We included do-not-resuscitate patients who received NIV in this study. We excluded the clinical data of those who refused informed consent.

Each Ethical Committee from all participating hospitals approved this study, and the Institutional Review Board of Hallym University also approved this study (IRB No. 2017-I049). Informed consent was obtained from all enrolled patients or their legal surrogates. We also performed a secondary analysis of NIV data in Korean ICUs. We compared the baseline characteristics and clinical data of patients receiving NIV with or without the use of pain and sedative drugs.

Data Collection and Outcomes

We collected demographic characteristics and the following data: comorbidities, presence and type of any underlying lung diseases, the primary causes of ICU admission and NIV treatment, mental status (the Richmond Agitation Sedation Score [RASS]), and the severity of the illness (the Sequential Organ Failure Assessment [SOFA]) immediately prior to the implementation of NIV. We also included the following in our analysis: arterial blood gas analysis (ABGA), vital signs, lactate level before and 2 hours after NIV, and the location where NIV was first implemented. While we collected the name of the
pain control and sedation drugs during NIV; we did not obtain information on the dose or duration. We also collected the specific type of NIV machine used, the NIV settings (i.e., main NIV modes, fractional inspired oxygen [FiO\(_2\)], inspiratory positive airway pressure [IPAP], expiratory positive airway pressure [EPAP], and tidal volume), and the mean duration of use (hr/day).

The following patient outcomes were examined: treatment success, treatment failure, complications from NIV treatment, length of NIV use, length of ICU stay, and ICU and hospital mortality rates. Treatment success was defined as: (1) successful weaning from NIV or (2) transfer from the ICU to a general ward in a stable state with the NIV device in place [15]. Treatment failure was defined as: (1) intubation and mechanical ventilation (MV), (2) tracheostomy, (3) hopeless discharge with a NIV device, (4) death during NIV treatment, or (5) change to the high-flow nasal cannula with no clinical improvement [16]. Patients who died within 24 hours of NIV weaning were also classified as a NIV failure [17]. A large leak was defined as a leak flow rate > 60 L/min, or when the attending doctor considered the leak too large to continue treatment.

Statistical Analysis
In this observational study, most statistical analyses were descriptive as the authors aimed to observe the current status of NIV use. All categorical variables are presented as numbers with percentages and were compared using the chi-square or Fisher’s exact test as indicated. All continuous variables are summarized and reported as means with standard deviations or medians with interquartile ranges. The normality test was conducted using the Shapiro-wilk test. The Student t-test was used if the variables had a normal distribution and the Mann-Whitney U-test was used if the variables had a skewed distribution. A paired t-test and the Wilcoxon signed-rank test were also used to compare the values before and after NIV use. Statistical analysis was performed using IBM SPSS ver. 22.0 (IBM Corp., Armonk, NY, USA) and the R statistical package version 4.0.0 (R Foundation, Vienna, Austria).

RESULTS
Baseline Characteristics
The baseline demographics of each group are summarized in Table 1. During the study period, a total of 155 patients treated with NIV were enrolled from 20 ICUs. This included 129 patients (83.2%) who did not undergo pain or sedation treat-

| Variable                          | Control (n = 129) | Sedation (n = 26) | P-value |
|----------------------------------|------------------|------------------|---------|
| Age (yr)                         | 71.7 ± 11.5      | 71.7 ± 11.4      | >0.05   |
| Male sex                         | 78 (60.5)        | 17 (65.4)        | >0.05   |
| SOFA score at NIV start          | 4.16 ± 0.24      | 4.27 ± 0.46      | >0.05   |
| RASS at NIV start                | −0.53 ± 1.06     | −0.11 ± 1.11     | 0.17    |
| Comorbidity                      |                  |                  | >0.05   |
| DM                               | 37 (27.1)        | 2 (7.7)          | 0.02    |
| HTN                              | 51 (39.5)        | 11 (42.3)        |         |
| Prior lung resection             | 6 (4.7)          | 2 (7.7)          |         |
| Coronary artery disease          | 16 (12.4)        | 4 (15.4)         |         |
| Heart failure                    | 29 (22.5)        | 5 (19.2)         |         |
| CKD                              | 17 (13.2)        | 3 (11.5)         |         |
| LC                               | 6 (4.7)          | 1 (3.8)          |         |
| CVA                              | 15 (11.6)        | 5 (19.2)         |         |
| Cancer                           | 10 (7.8)         | 3 (11.5)         |         |
| Immunocompromised state          | 8 (6.2)          | 1 (3.8)          |         |
| Underlying lung disease          |                  |                  | >0.05   |
| Normal                           | 35 (27.1)        | 6 (23.1)         |         |
| COPD                             | 44 (34.1)        | 10 (38.5)        |         |
| Bronchial asthma                 | 3 (2.3)          | 0                |         |
| ILD                              | 4 (3.1)          | 2 (7.7)          |         |
| Bilateral bronchiectasis         | 7 (5.4)          | 3 (11.5)         |         |
| TB-destroyed lung                | 15 (11.6)        | 3 (11.5)         |         |
| Neuromuscular                    | 6 (4.7)          | 0                |         |
| Obesity disorder                 | 4 (3.1)          | 1 (3.8)          |         |
| Kyphoscoliosis                   | 4 (3.1)          | 0                |         |
| Other chest wall disease         | 2 (1.6)          | 0                |         |
| Obstructive sleep apnea          | 3 (2.3)          | 0                |         |
| Others                           | 2 (1.6)          | 1 (3.8)          |         |

Values are presented as mean ± standard deviation or number (%).

Table 1. Baseline demographics and characteristics of the study population

Statistical analysis was performed using IBM SPSS ver. 22.0 (IBM Corp., Armonk, NY, USA) and the R statistical package version 4.0.0 (R Foundation, Vienna, Austria).


dent (control group) and 26 patients (16.8%) who underwent analgesedation during NIV (sedation group) (Figure 1). The mean age of all the enrolled patients was 71.7 ± 11.6 years, of whom 95 patients (61.3%) were male. At the start of NIV, the mean SOFA score was 4.16 ± 0.24 in the control group and 4.27 ± 0.46 in the sedation group (P > 0.05). The RASS was −0.53 ± 1.06 in the control group and −0.11 ± 1.11 in the sedation group for mentality evaluation before NIV was applied.
Hypertension was the most common comorbidity in both groups: 39.5% (51/129) in the control group and 42.3% (11/26) in the sedation group (P = 0.02). COPD as an underlying lung disease was reported 34.1% (44/129) in the control group and 38.5% (10/26) in the sedation group. However, the most common cause of ICU admission in the sedation group was pneumonia treatment (14/26, 53.8%) (Table 2). In addition, the most common causes of NIV application were acute hypercapnic respiratory failure in the control group (62.8% [81/129]) and post-extubation respiratory failure in the sedation group (57.7% [15/26]).

Changes in Vital Signs and ABGA

We compared alterations in vital signs and ABGA results between baseline values before NIV treatment and 2 hours after NIV treatment in both groups (Table 3). In the control group, arterial partial pressure of carbon dioxide (PaCO₂) was significantly reduced from 61.9 ± 23.8 mm Hg before NIV treatment to 54.9 ± 17.6 mm Hg after 2 hours of NIV treatment (P < 0.001). Heart rate (HR) was 95.5 ± 19.7 before NIV and 90.7 ± 16.1 after 2 hours of NIV treatment (P < 0.001). The respiratory rate (RR)
was 24.5 ± 6.72 before NIV and 23.4 ± 5.87 after 2 hours of NIV (P = 0.018). However, pH, ratio of arterial oxygen partial pressure (PaO₂), and systolic blood pressure (SBP) did not exhibit interval changes before and after NIV treatment in the control group. In the sedation group, PaCO₂ was significantly reduced from 54.9 ± 15.1 mm Hg before NIV treatment to 51.1 ± 15.1 mm Hg after 2 hours of NIV treatment (P = 0.048). SBP was reduced from 139 ± 25.1 mm Hg before NIV to 127 ± 19.6 mm Hg after 2 hours of NIV treatment (P = 0.021). There were no significant differences in pH, PF ratio, HR, and RR in the sedation group.

**NIV Treatments (Devices, Modes, and Settings)**

The differences in the type of NIV machine and the applied NIV mode were analyzed in both the control and sedation groups. In the control group without sedation, an ICU ventilator with an NIV mode was used for 73 cases (56.6%), a dedicated NIV was used for 29 cases (22.5%), and a home care NIV unit was used for 25 cases (19.4%). In the sedation group, an ICU ventilator with an NIV mode was used for 29 cases (22.5%), and a dedicated NIV was used for 23 cases (88.5%), and a dedicated NIV was used for three cases (11.5%, P = 0.009). In the control group, the pressure support ventilation (PSV) mode was used in 56 patients (84.8%). In the sedation group, the pressure control ventilation mode was used in 13 patients (50%) and the PSV mode was used in 10 patients (38.5%) (P = 0.016).

The mean IPAP was 14.5 ± 0.36 cm H₂O in the control group and 14.1 ± 0.49 cm H₂O in the sedation group (P = 0.386), and the mean EPAP was 5.05 ± 0.12 cm H₂O and 4.92 ± 0.37 cm H₂O (P = 0.229), respectively. The tidal volume was 432 ± 16.1 ml in the control group and 529 ± 51.0 ml in the sedation group (P = 0.05).

**Clinical Outcomes and Complications**

Forty-six patients (35.7%) in the control group and 14 patients (53.8%) in the sedation group had failed NIV treatment (P = 0.086). In the control group, 22 patients (17.1%) underwent tracheal intubation and maintained MV. In the sedation group, tracheal intubation was performed in eight patients (30.8%). The most common cause of NIV failure in both groups was inadequate efficacy (i.e., no clinical improvement or no improvement in ABGA). In the sedation group, three cases (21.4%) had failed NIV due to copious secretions. The length of ICU stay was 16.1 ± 1.5 days in the control group and 17.8 ± 2.80 days in the sedation group (P = 0.210). The length of the NIV period was 4.78 ± 0.39 days in the control group and 5.28 ± 0.38 days in the sedation group (P = 0.001). In the sedation group, the length of the NIV period was significantly shorter than that of the control group. However, this was due to the NIV weaning failure being more frequent in the sedation group (Table 4).

**Table 4. Clinical outcomes after NIV between the two groups**

| Outcome                          | Control (n = 129) | Sedation (n = 26) | P-value |
|---------------------------------|------------------|------------------|---------|
| NIV failure                     | 46 (35.7)        | 14 (53.8)        | 0.086   |
| Intubation and MV               | 22 (17.1)        | 8 (30.8)         |         |
| Tracheostomy                    | 4 (3.1)          | 3 (11.5)         |         |
| Hopeless discharge with NIV     | 3 (2.3)          | 0                |         |
| Died during NIV                 | 13 (10.1)        | 2 (7.7)          |         |
| Change to HFNC                  | 11 (3.1)         | 3 (11.5)         |         |

| Cause of NIV failure            | 0.021            |                   |         |
|                                 |                  |                   |         |
| Inadequate efficacy             | 36 (80.0)        | 8 (57.1)          |         |
| No clinical improvement         | 22 (47.8)        | 4 (28.6)          |         |
| Lack of arterial blood gas      | 14 (30.4)        | 4 (28.6)          |         |
| improvement                     |                  |                   |         |
| Copious secretion               | 0                | 3 (21.4)          |         |
| Agitation                       | 1 (2.2)          | 1 (7.1)           |         |
| Large leaks                     | 2 (4.4)          | 0                |         |
| Patients’ discomfort or refuse  | 6 (13.3)         | 2 (14.3)         |         |
| Length of NIV (day)             | 4.78 ± 0.39      | 2.27 ± 0.38       | 0.001   |
| Length of ICU stay (day)        | 16.1 ± 1.50      | 17.8 ± 2.80       | 0.210   |
| ICU mortality                   | 19 (15.0)        | 3 (11.5)          | 0.769   |
| Hospital mortality              | 28 (22.0)        | 7 (26.9)          | 0.590   |

Values are presented as number (%) or mean ± standard deviation. NIV: noninvasive ventilation; MV: mechanical ventilation; HFNC: high-flow nasal cannula; ICU: intensive care unit.

**Table 5. Safety outcomes during NIV between the two groups**

| Outcome                                         | Control (n = 129) | Sedation (n = 26) | P-value* |
|-------------------------------------------------|------------------|------------------|----------|
| No. of complications                            | 45 (34.9)        | 9 (34.6)         | 0.979    |
| Complication during NIV                         |                  |                  |          |
| Skin erythema                                    | 20 (15.5)        | 6 (23.1)         | 0.389    |
| Abdominal distension                            | 6 (4.7)          | 1 (3.8)          | 1.000    |
| Dry mouth                                        | 5 (3.9)          | 1 (3.8)          | 1.000    |
| Aspiration                                       | 5 (3.9)          | 0                | 0.590    |
| Large leak                                       | 16 (12.4)        | 1 (3.8)          | 0.309    |
| Claustrophobia                                   | 4 (3.1)          | 0                | 1.000    |
| Nasal congestion or sinus pain                  | 0                | 1 (3.8)          | 0.168    |
| Mucosal plugging                                | 1 (0.8)          | 0                | 1.000    |

Values are presented as number (%). NIV: noninvasive ventilation. *Fisher exact test.
Complications of NIV were reported in 45 patients (34.9%) in the control group and nine patients (34.6%) in the sedation group. The most commonly reported complication during the NIV period was skin erythema, with 20 cases (15.5%) occurring in the control group and six (23.1%) in the sedation group (Table 5).

**Pharmacologic Use and Type**

Of all the patients, 26 patients (16.7%) received a sedative (or analgesic) during NIV treatment with remifentanil (n = 15) and dexmedetomidine (n = 8) being the most commonly used (Table 6).

| Medication     | Sedation group (n = 26) |
|----------------|-------------------------|
| Remifentanil   | 15 (57.7)               |
| Dexmedetomidine| 8 (30.8)                |
| Fentanyl       | 1 (3.8)                 |
| Midazolam      | 1 (3.8)                 |
| Morphine       | 1 (3.8)                 |

Values are presented as number (%). NIV: noninvasive ventilation.

Not a significant difference. Skin erythema was the most common complication of NIV in both groups.

**DISCUSSION**

Our study contains several important findings. First, we found that 16.8% of patients with NIV treatment received pain or sedation therapy in Korean ICUs. In the sedation group, the most common cause for the implementation of NIV was post-extubation respiratory failure (57.7%). In the same group, the most common cause of ICU admission was pneumonia treatment (53.8%). Second, while the rate of NIV failure was 35.7% in the control group and 53.8% in the sedation group, the difference was not statistically significant. Furthermore, the rate of tracheal intubation was 17.1% in the control group and 30.8% in the sedation group. NIV was primarily applied to the sedation group post-extubation, which may be the result of extubation failure. This hypothesis is supported by the observation that a portion of the NIV failures in the sedation group were due to copious secretions (21.4%). Third, the length of NIV use was 2.5 days shorter in the sedation group when compared to the control group. The NIV failure rate in the sedation group was high, which may explain this shorter NIV duration. However, there were no differences in the length of ICU stay, ICU mortality, or hospital mortality between the two groups. Finally, the incidence of NIV complications was 34.9% in the control group and 34.6% in the sedation group, which is not a significant difference. Skin erythema was the most common complication of NIV in both groups.

There are several limitations in this study. First, the number of patients who underwent sedation in our total patient population was insufficient. Devlin et al. [18] conducted a survey of 2,985 physicians in North America and Europe and found that 41% used sedative drugs in NIV in North America and 24% in Europe. However, only 17% of patients in Korean ICUs who underwent NIV used sedative drugs. The authors [18] reported that the regional differences in the use of sedative drugs in NIV treated patients was due to differences in physician experience in applying NIV, experience in sedation monitoring, presence of a sedation protocol, and the nurse-to-patient ratio. In a survey by Devlin et al. [18], the most widely used classes of sedation drugs were benzodiazepines (33%) and opioids (29%), and the most influential factors determining the choice of sedative were clinical experience with the agent (33%) and a lack of an effect on respiratory drive (22%). In our study, the most commonly used sedative drugs during NIV were remifentanil (57.7% [15/26]) and dexmedetomidine (30.8% [8/26]). However, the reasons for choosing these specific pharmacologies and the most efficacious sedative drug dose were not investigated. Therefore, further investigation and analysis of the most efficacious sedative drug doses and the specific reasons for the selection of the drugs is needed in future studies. Third, the reasons for choosing NIV treatment between the control and the sedation groups were different in our study. It is well-known that the most common indications for NIV are the acute exacerbation of COPD and cardiogenic pulmonary edema. In our study, the primary reason for NIV was hypercarbic respiratory failure (63.6%) in the control group and post-extubation for weaning (57.7%) in the sedation group. This difference may be due to the required maintenance of sedation during NIV after extubation in patients who used sedative drugs during MV treatment. In addition, NIV after extubation is clinically more likely to result in lower voluntary respiration weaning rates and higher rates of re-intubation. This could then explain the higher NIV failure rate in the sedation group. It also suggests that clinicians should be intensive in their observations of patients during the use of sedation. Despite these limitations, the present study provides the first extensive survey of current sedation practices in patients treated with NIV in Korean ICUs.

In conclusion, for patients who were admitted to Korean ICUs and treated with NIV, the use of sedatives did not present any obvious harmful effects in regard to ICU hospital stay, ICU mortality or hospital mortality, or NIV complications.
Therefore, we suggest the application of analgosedation may not be harmful for patients who are undergoing NIV.

CONFLICT OF INTEREST

Jae Hwa Cho has been the editor-in-chief of Acute and Critical Care since 2016 and an editorial board member since 2008.

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Conceptualization: TK, JSK, EYC, SP, JHC. Data curation: all authors. Formal analysis & Methodology: TK, JSK, SP, JHC. Visualization: TK, JHC. Writing–original draft: TK, SP, JHC. Writing–reviewing & editing: all authors.

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