Original Research Article

Effectiveness of non-invasive ventilation in patients with type 2 respiratory failure

Shahid M. Patel*, Girija P. Nair, Balaji G. Tuppekar, Abhay G. Uppe,

Department of Pulmonary Medicine, D Y Patil Medical college, Navi Mumbai, Maharashtra, India

Received: 03 January 2021
Revised: 05 February 2021
Accepted: 06 February 2021

*Correspondence:
Dr. Shahid M. Patel,
E-mail: patel_shahid@hotmail.com

ABSTRACT

Background: Assess the use of non-invasive ventilation as an alternative way for ventilation in acute respiratory failure, determine factors that can predict the successful use of NIV, evaluate factors hindering success of NIV.

Methods: Thirty hospitalised patients fulfilling inclusion criteria, diagnosed with Type II Respiratory Failure on ABG were recruited after obtaining an informed written consent. Complete history and detailed physical examination were followed by routine investigations.

Results: Comparison of the pH on admission with the pH after 1st hour of NIV, the latter showed statistically significant improvement. Drop in PaCO2 and rise in PaO2 on ABG from admission and after stopping NIV was statistically significant. Patients with lower MMRC grade and severe cough showed significant improvement in pH, however patient with higher emergency visits and past hospitalisation showed less improvement in pH, after 1 hour of NIV therapy. A total 4 patients were intubated, with mean pH of 7.22, 3 out of them had higher emergency visits, 2 out of them had ICU admission.

Conclusions: NIV treatment for COPD with type II respiratory failure avoids intubation, reduces complications and should be considered as first line therapy instead of ET intubation. Lower mMRC grade, lesser hospitalizations, lesser emergency visits, higher BMI, symptoms like cough, can have a positive predictive value for the outcome of NIV.

Keywords: Non-invasive ventilation, Acute respiratory failure, ABG, COPD

INTRODUCTION

NIV is one of the treatment modalities for type 2 respiratory failure. NIV has been used successfully to treat acute respiratory failure in postoperative patients, in those with pulmonary edema, COPD and obstructive sleep apnea. It has also been used to facilitate weaning. However, NIV appears to be particularly effective in patients with an exacerbation of COPD, who are alert and cooperative. Patel et al found that respiratory rate significantly improves during the first 30 minutes after the application of NIV treatment. Although, many researchers have studied NIV, still there is a paucity of literature.

With limited resources, the modality of the treatment has to be of greater relevance.

Thus, this study was done with the aim to assess the use of non-invasive ventilation as an alternative way for ventilation in acute respiratory failure, determine factors that can predict the successful use of NIV, evaluate factors hindering success of NIV.
METHODS

It was a prospective longitudinal study conducted on thirty hospitalized patients fulfilling inclusion criteria. The study was conducted from January 2018 to September 2020 at D Y Patil Hospital, Nerul, Navimumbai. Patients diagnosed with Type II Respiratory Failure on ABG, having symptoms of breathlessness, cough, use of accessory muscles of respiration, associated findings of increased PaCO2 like malaise, drowsiness, asterexis, anxiety, restlessness, confusion were recruited after obtaining a written informed consent.

Complete history and detailed physical examination were followed by routine investigations like chest x-ray, ECG, complete blood count, renal function test, liver function test, PFT, 2D ECHO and CT scan of the chest.

The patients were started on NIV therapy, along with the standard symptomatic medical therapy as appropriate. ABG was done on admission and after 1 hour.

Success was considered when the pH returned to normal, along with normalizing of the Patients physical and biochemical parameters, i.e., normalizing of the pH on ABG, reduction in the Respiratory rate, reduction in the Pulse rate, and decrease in the patient’s sensation of Dyspnœa. Patients were put on a Biphasic/Bilevel ventilation (BIPAP) machine with an initial setting of IPAP of 12 cm of H2O and an EPAP of 6 cm of H2O. The setting was stepped up, in required cases.

Data was collected in MS-Excel and all statistical analysis was done in statistical package for social sciences (SPSS) -27. Categorical data was analysed using chi-square test and continuous data was analysed using t-test. Significance was checked at 95% confidence interval (p<0.05). The study was approved by ethical committee.

Indications of NIV

Acute hypercapnic exacerbations of COPD in preventing intubation of end-stage COPD patient. OSA: caused by airflow obstruction, is defined as a temporary pause in breathing that lasts at least 10 sec during sleep. Central sleep apnea - loss of neurologic breathing effort (CSA), or a combination of both mixed sleep apnea (MSA). Reduction of respiratory workload in obesity. Acute pulmonary edema.

Contraindications for use of NIV

Apnoea due to neuromuscular causes and progressive hypoventilation. Inability to handle secretions by the patient. Hypotension and fatigue of respiratory muscles. Claustrophobia and facial trauma.

Inclusion criteria

All patients of age 18 years or greater after consent, indoor patients of Padmashree Dr. D.Y. Patil Hospital and Research Centre diagnosed as being in type ii respiratory failure after arterial blood gas studies.

Exclusion criteria

Patients with facial surgery, trauma or deformity, cardiac or respiratory arrest were excluded. Also, those with presence of upper airway obstruction, hypotension (systolic BP <90 mmHg), uncontrolled arrhythmia, severe upper gastrointestinal bleeding, not able to clear respiratory secretions with high risk for aspiration, and other non-respiratory organ failure: example: Severe encephalopathy (example: GCS <10) were also excluded.

RESULTS

The aim of this study was to evaluate various factors for predicting the success or failure of a NIV trial. The mean age was 59.9±10.13 with a Male pre-dominance, 3:2. The mean BMI was 22.9±3.97 kg/m2. Out of 30 patients, 13 were smokers, 10 were cigarette smokers and 3 were bidi smokers.

![Figure 1: Distribution based on age group.](image1)

![Figure 2: Distribution based on age group and smokers.](image2)

The most common comorbidities in our patients were hypertension, diabetes mellitus and ischemic heart disease. The pH improvement was more in patients who did not have these comorbidities.
Fourteen (46.6%) of the patients had dyspnea on exertion Grade 2 MMRC, 11 (36.6%) had MMRC Grade 1 and less, 5 (16.6%) had grade 3. The pH improvement was better in patients who presented with low MMRC grade. Out of 30, 28 patients had cough and the pH improvement had a positive correlation with duration of cough. Eighty-eight percentage (n=16) of patients had fever and the pH improvement had a negative correlation with fever.

Out of the 30 patients, all survived, and 4 had to be intubated, and hence were considered as failure of the NIV trial. The mean pH difference after 1 hour of NIV in the 26 patients was +0.07, while the mean pH difference after 1 hr in the patients who had to be intubated was -0.0005.

Prior studies have shown that the change in pH in the initial period of the NIV trial, is an indicator for the outcome of the trial, i.e. higher the change, it’s more likely that the NIV trial will succeed. Hence for the purpose of our study, the change in the value of pH after 1 hour, was considered as an outcome, and the various factors were statistically analysed with the change in pH after 1 hour and results observed were that factors such as compliance with medications, lower number of hospitalizations in last 5 years, lower number of emergency visits in last 6 months, lower mMRC grade prior to exacerbation showed more change in pH in the first hour as compared to otherwise.

Twenty-eight patients had cough out of which 21 had expectoration, 16 had complaints of fever and 29 patients had complaints of Dyspnoea for some duration. While the duration of fever and duration of dyspnoea were found to be inversely proportional to the improvement in the pH in the first hour, patients with longer duration of cough showed more improvement in pH. This might be attributed to the fact that patients with a longer duration of cough, might have taken medications prior to admission. The smoking duration and the chulla exposure duration were found to have an inverse relation to the change in the pH in the first hour. A lower BMI was found to have a lower change in the pH after 1 hour, this is concurring with a study published in BMC Pulmonary medicine in 2014. Our study also showed that, when comparing the pH on admission with the pH at 1st hour and pH on withdrawal of NIV, the pH after 1 hour was statistically significant (p<0.0001), which is consistent with other studies on NIV, but when compared to the pH at withdrawal it was found to be statistically not relevant.

The PaCO2 after 1 hour and at withdrawal, both were found to be statistically significant (p value<0.0001) when compared to the PaCO2 on admission. This study shows that factors such as presence of co-morbidities, MMRC grade prior to the exacerbation episode, compliance of patients with medications, history of hospitalizations in last 5 years, history of emergency visits, duration of dyspnea, smoking history, history of chulla exposure, can be considered a prognostic factor for the success of the NIV trial. The profile of the NIV failure cases, which had to be intubated have been mentioned in table 1, the mean pH was 7.22, with the mean duration of NIV trial was 13.5 hours, which is consistent with other studies showing that the majority of NIV failure occur within 48 hours. Out of the 4 patients, 3 had a higher number of emergency visits and a higher no hospitalizations, with 2 having Intensive Care Unit admissions and also all had some findings on the 2D-echo.

**DISCUSSION**

The mean duration of BiPAP use was 4.03±2.11 days. Out of the 30, 10 were cigarette smokers and 3 were bidi smokers out of which 4 were still active in the cigarette smoking group and 1 was active in the bidi smoking group. Six had been exposed to chulla smoke, out of which 2 still had continued exposure.
In our study we found that the maximum pH improvement in the first hour was seen in the case of the patient with the diagnosis of ILD. She was a 60/F, with 20 years of chulla exposure, a good BMI, with no co-morbidities and had no emergency visits and only one incidence of hospitalization, and these factors could be the reason behind her better improvement. She had come to us with a HRCT Chest showing ILD changes but her history and symptoms were more consistent with COPD. The pH improvement was more in Obesity-Hypoventilation group, followed by acute exacerbation of COPD then others. The poorest improvement was seen in the infective exacerbation of COPD group. In obesity-hypoventilation, there is no lung pathology as such, and hence NIV helps in improving the ventilation.13

The minimum duration of use of the NIV was found to be in the acute exacerbation of COPD group, while the others had approximately the same mean duration of use.14 When comparing acute exacerbation of COPD versus infective exacerbation of COPD, the first group had a greater improvement in the pH after 1 hour and also the mean duration of the NIV was lesser (almost half as that of the infective group). This could be because the infective group was more likely to have a more severe illness.15

The main limitation of the study was that it was a single centre study. Another limitation of this study was its small sample size and lack of control group. Due to which there was limited value to statistical analysis. So, the outcomes may seem inconclusive.

CONCLUSION

ABG values after 1 hour of institution of NIV therapy, is very significant and can be used as a predictor of outcome of NIV. Lower mMRC grade, lesser hospitalizations, lesser emergency visits, higher BMI, also have a positive predictive value for NIV outcomes. NIV treatment for COPD with type II respiratory failure avoids intubation, reduces complications and should be considered as first line therapy instead of ET intubation. By avoiding endotracheal intubation, NIV prevents complications associated with invasive ventilation like airway problems, nosocomial pneumonia (21%) and sinusitis (5-25%).

Funding: No funding sources
Conflict of interest: None declared
Ethical approval: The study was approved by the Institutional Ethics Committee

REFERENCES

10. Ambrosino N, Vagheggini G. Noninvasive positive pressure ventilation in the acute care setting: where are we? Eur Respir J. 2008;31:874-86.