Comparison of the acute effects on gas exchange of nasal ventilation and doxapram in exacerbations of chronic obstructive pulmonary disease

R M Angus, A A Ahmed, L J Fenwick, A J Peacock

Abstract

**Background** – Nasal intermittent positive pressure ventilation (NIPPV) is useful in exacerbations of chronic obstructive pulmonary disease (COPD) complicated by ventilatory failure. The effects of NIPPV were compared with those of the respiratory stimulant doxapram on gas exchange in patients with COPD and acute ventilatory failure. **Methods** – Patients admitted with acute exacerbations of COPD and type 2 respiratory failure (Pao2 <8 kPa and PacO2 >6.7 kPa) who did not improve with conventional treatment were randomised to receive either NIPPV or intravenous doxapram. Blood gas tensions were monitored for four hours. **Results** – In nine patients who received NIPPV the arterial Pao2 improved from a mean (SE) of 5.9 (0.4) kPa to a maximum of 8.1 (0.6) kPa which was maintained at four hours. Eight patients who received doxapram had a similar baseline Pao2 of 5.6 (0.4) kPa which rose to a maximum of 7.3 (0.5) kPa but this was not maintained at four hours. The improvement in Pao2 in patients on NIPPV was accompanied by a fall in PacO2, but, in contrast, in those who received doxapram there was no improvement in PacO2. **Conclusions** – NIPPV may be more effective than doxapram in the management of acute ventilatory failure complicating COPD. (Thorax 1996;51:1048–1050)

Keywords: respiratory failure, NIPPV, doxapram.

The usual cause of death in patients with chronic obstructive pulmonary disease (COPD) is respiratory failure. Until recently the only readily available forms of ventilatory support were intravenous doxapram or mechanical ventilation via an endotracheal tube. Doxapram drives respiration, probably by stimulating peripheral chemoreceptors. This is helpful if there is suppression of central drive but, in the context of an acute exacerbation of chronic lung disease, the respiratory muscles may already be functioning at near maximal capacity. Intubation and full scale intermittent positive pressure ventilation is invasive, expensive, and it may be difficult to wean the patients off the ventilator. An effective alternative would be desirable.

Nasal intermittent positive pressure ventilation (NIPPV) has also been shown to improve arterial blood gas tensions in patients with COPD, but no comparison has been published of the effects of NIPPV and doxapram in the emergency treatment of ventilatory failure. We have therefore compared NIPPV and doxapram in a group of patients presenting to a respiratory ward with type 2 respiratory failure.

**Methods**

**Patients**

Patients were included if they had an acute exacerbation of COPD complicated by type 2 respiratory failure (Pao2 <8 kPa and PacO2 >6.7 kPa). Seventeen patients completed the study; nine (five men) of mean (SE) age 64 (2.3) years and forced expiratory volume in one second (FEV1) 0.76 (0.02) 1 received NIPPV and eight (four men) of mean (SE) age 61.9 (2.7) years and FEV1 0.77 (0.09) 1 received doxapram. The study was approved by the Glasgow West ethical committee and informed consent was obtained.

**Study Design**

Blood gas tensions were measured on admission. Patients were then started on routine treatment with oxygen (1–3 l/min by nasal cannulae), steroids, antibiotics, and nebulised or intravenous bronchodilators. One hour later arterial blood gas tensions were repeated. If the values were static or deteriorating, the patient was randomised to receive either nasal ventilation or doxapram. Oxygen was continued at the same rate as before randomisation. NIPPV was administered via a pressure cycled machine (Ventimate, Thomas Respiratory Systems, London, UK). Ventilation pressure was set at...
Nasal and doxapram in patients

![Graph](image)

Figure 1 Mean (SE) changes in PaO2 and PaCO2 in patients admitted with acute respiratory failure treated with either intravenous doxapram (n=8) or nasal intermittent positive pressure ventilation (NIPPV, n=9).

* p<0.05 compared with baseline.

14–18 cm H2O and oxygen enriched air was delivered using nasal pillows or a tight fitting nasal mask. Doxapram was given by intravenous infusion using the manufacturer's protocol (4 mg/min for 15 minutes, 3 mg/min for 30 minutes, 2 mg/min for 60 minutes, and 1.5 mg/min as maintenance dose). Arterial blood gas tensions were checked hourly for the first four hours. The protocol was modified later so that if there was continued deterioration on either NIPPV or doxapram the alternative was added at four hours.

DATA ANALYSIS

Arterial blood gas tensions were compared using ANOVA. pH was converted to H+ concentration, compared by ANOVA, and reconverted to pH expressed as the geometric mean + 95% confidence intervals. p values of less than 0.05 were regarded as significant.

Results

At entry the baseline blood gas tensions were similar in the two groups. The changes in arterial blood gas tensions during the four hour assessment period are shown in fig 1. The mean (SE) baseline PaO2 of 5.9 (0.4) kPa improved on NIPPV to a maximum value of 8.1 (0.6) kPa (p<0.05) which was maintained at four hours. The baseline PacO2 of 10.1 (0.4) kPa also improved on NIPPV to 8.7 (0.6) kPa (p<0.05), and the improvement was maintained at four hours.

In patients receiving doxapram the mean (SE) baseline PaO2 of 5.6 (0.4) kPa improved on treatment to a maximum value of 7.3 (0.5) kPa (p<0.05), but this was not maintained at four hours. The baseline PacO2 of 10.1 (0.8) kPa did not improve on treatment. At four hours the change in PaO2 was significantly greater in the NIPPV group than in the doxapram group.

Discussion

Neither group was severely acidotic (mean pH 7.30 in the doxapram group and 7.31 in the NIPPV group). Changes in pH during the study are shown in fig 2.

All patients tolerated NIPPV but doxapram was added in one because of drowsiness at four hours. All patients who received NIPPV were discharged home.

Three patients in the doxapram group died after four hours but within the first 24 hours. During the first four hours their gas tensions had continued to deteriorate despite treatment and a decision was made not to offer full mechanical ventilation. After these three deaths the protocol was altered to allow the addition of the alternative form of respiratory support. Two further patients on doxapram had NIPPV added at four hours because their clinical condition and blood gas tensions were deteriorating (PaCO2 rose from 8.5 to 8.7 and pH fell from 7.31 to 7.28). Their arterial blood gas tensions improved on NIPPV (mean PaO2 improved from 7.6 kPa to 11.3 kPa and mean PacO2 improved from 8.7 kPa to 8.3 kPa). They both survived to discharge.

This study has shown that non-invasive nasal ventilation produces an immediate improvement in gas exchange in patients with ventilatory failure complicating an exacerbation of COPD, and that this improvement is maintained for at least four hours. It is possible that the fractional inspired oxygen (FiO2) was altered by the insertion of the nasal mask circuit but the fall in PacO2 with NIPPV suggests an improvement in ventilation. In contrast, on doxapram an initial improvement in PaO2 was not maintained. There are three explanations for this late deterioration. Firstly, the prescribing guidelines recommending a high initial loading dose followed by a reduction may be inappropriate. Secondly, although matched and randomised, the doxapram group may have simply...
deteriorated more rapidly. Thirdly, and most likely, with the respiratory muscles already working at near capacity a stimulant may offer only temporary benefit until fatigue supervenes. In contrast, assisted ventilation may offload the respiratory muscles.

Between four and 24 hours three patients in the doxapram group died. Two further patients who deteriorated in the doxapram group were treated successfully by the addition of NIPPV. One patient in the NIPPV group received doxapram, resulting in better cooperation with the nasal ventilation. While it might have been possible to alter ventilator settings rather than add doxapram, we would suggest that the addition of doxapram might improve acceptance of NIPPV, particularly if, as here, the patient was drowsy.

Despite the lack of evidence for successful treatment of ventilatory failure using doxapram in patients with acute infective exacerbations of COPD, it is widely used for this purpose. In contrast, there is good evidence for the effectiveness of NIPPV in this situation.4,6 We have now shown in a small study that NIPPV is more effective than doxapram in these patients. We suggest that doxapram is reserved for patients who are centrally sedated or are unable to tolerate nasal ventilation.

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