A Postoperative Ventilatory Failure

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Abstract—A case of severe ventilatory failure with PaCO₂, as high as 197 torr is reported postoperatively in a 3-year-old male patient who has received open heart surgery for total correction of known TOF (Tetralogy of Fallot). Many procedures were tried to treat the ventilatory failure but failed. Eventually we noticed malfunctioning of the humidifier assembled in Bear Cub infant pressure-limited ventilator. Vigorous endotracheal suction combined with saline irrigation improved the ventilatory failure successfully.

Key Words: Ventilatory insufficiency, TOF, Humidifier, Respiratory care

Inadequate physiologic ventilation would be an arterial PCO₂ above an acceptable range which is a condition referred to as ventilatory failure or respiratory acidosis.

Assessment of the adequacy of physiologic ventilation can be accomplished only by blood gas analysis (Shapiro et al. 1982). Inadequate alveolar ventilation (PaCO₂ above an acceptable limit) reflects a failure to remove CO₂ adequately through the lungs. When such a "ventilatory failure" is accompanied by acidemia (arterial pH below an acceptable limit), the failure must be of recent origin (acute) and, therefore, a direct threat to cardiopulmonary homeostasis. Acute ventilatory failure may be defined arbitrarily as PaCO₂ above 50 torr coincidental with an arterial pH below 7.30 (Shapiro et al. 1985). Recently we experienced a case of severe acute ventilatory failure during mechanical ventilatory support, which was treated successfully with vigorous endotracheal suction combined with saline irrigation.

CASE REPORT

A 3-year-old male patient underwent an open heart surgery of TOF under general anesthesia composed of N₂O-O₂-Halothane with intermittent intravenous injection of morphine. After successful operation, he was transferred to RICU (Respiratory Intensive Care Unit) in intubated state and connected to Bourns infant volume ventilator (FiO₂ 0.7, tidal volume 110 ml, respiratory rate 25/min.). ABGA (Arterial Blood Gas Analysis) was within normal limit. On the postoperative 2nd day, bilateral pulmonary edema and pleural effusion were detected by a chest roentgenogram (Fig. 1). Chest tubes were inserted. 5 cmH₂O of PEEP (Positive End-Expiratory Pressure) was applied. On the 6th day, ventilator was switched to Bear Cub infant ventilator because of danger of pulmonary barotrauma. Airway pressure was 35 cmH₂O under 12 cmH₂O of PEEP. Endotracheal pinkish and frothy secretion and pulmonary edema were still present. On the 11th day, ABGA showed pH 6.90, PaCO₂ 197 torr and PaO₂ 104 torr at FiO₂ 1.0, respiratory rate 60/min. and PEEP 5 cmH₂O. Airway pressure was 45 cmH₂O. Repeated attempts of endotracheal suction and manual ventilation with Ambu resuscitator were unsuccessful to correct the acidosis and ventilatory failure. A suction catheter could not be inserted beyond a certain point through trachea. A chest roentgenogram confirmed the diagnosis of bilateral pneumothorax, indicating pulmonary barotrauma (Fig. 2). On the postoperative 12th day, it was noted that the humidifier attached to Bear Cub infant ventilator was malfunctioning, and after several attempts of vigorous endotracheal suction and saline irrigation, aggregated blood clots and secretions were expelled. Thoracic movement got better and airway pressure was dropped from 50 cmH₂O to 40 cmH₂O immediately after then. Acute
Fig. 1. Diffuse bilateral alveolar consolidation is seen in both lung fields, suggesting pulmonary edema. Linear radiolucency in left upper portion of mediastinum suggests localized pneumomediastinum. Chest tubes are inserted in both sides.

Fig. 2. Extensive bilateral pneumothorax is seen, suggesting pulmonary barotrauma. Pulmonary edema shows no significant interval change compared with Fig. 1. Bilateral chest tubes are inserted.
Table 1. Postoperative changes of arterial blood gas

<table>
<thead>
<tr>
<th>POD</th>
<th>pH-PaCO₂-PaCO₂ (torr) (torr)</th>
<th>F₁O₂</th>
<th>V₁(ml)</th>
<th>t/min.</th>
<th>PEEP (cmH₂O)</th>
<th>AP (cmH₂O)</th>
<th>Ventilation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>7.41- 41- 214</td>
<td>0.7</td>
<td>110</td>
<td>25</td>
<td>5</td>
<td>NC</td>
<td>Bourns</td>
</tr>
<tr>
<td>2</td>
<td>7.53- 41- 67</td>
<td>0.5</td>
<td>110</td>
<td>25</td>
<td>5</td>
<td>NC</td>
<td>&quot;</td>
</tr>
<tr>
<td>5</td>
<td>7.48- 42- 69</td>
<td>0.7</td>
<td>130</td>
<td>30</td>
<td>8</td>
<td>NC</td>
<td>&quot;</td>
</tr>
<tr>
<td>6</td>
<td>7.39- 53- 54</td>
<td>1.0</td>
<td>130</td>
<td>40</td>
<td>12</td>
<td>NC</td>
<td>&quot;</td>
</tr>
<tr>
<td>11</td>
<td>7.38- 37- 83</td>
<td>1.0</td>
<td>NC</td>
<td>50</td>
<td>12</td>
<td>35</td>
<td>Bear</td>
</tr>
<tr>
<td>12</td>
<td>7.40- 51- 75</td>
<td>0.8</td>
<td>NC</td>
<td>60</td>
<td>5</td>
<td>40</td>
<td>&quot;</td>
</tr>
<tr>
<td>16</td>
<td>7.45- 38- 130</td>
<td>0.3</td>
<td>120</td>
<td>5</td>
<td>5</td>
<td>NC</td>
<td>&quot;</td>
</tr>
</tbody>
</table>

POD: Postoperative day  
AP: Airway pressure  
NC: Not checked  
f: Respiratory frequency  
F₁O₂: Fraction of inspired oxygen  
V₁: Tidal volume  

Ventilatory failure was treated completely. ABGA was pH 7.59, PaCO₂ 30 torr and PaO₂ 98 torr at F₁O₂ 0.5, respiratory frequency 40/min, and PEEP 5 cmH₂O in Bear Cub infant ventilator. The ventilator was switched to Bourns infant volume ventilator which was set at tidal volume 130 ml, frequency 30/min., PEEP 5 cmH₂O and F₁O₂ 0.5. In this situation, ABGA was pH 7.56, PaCO₂ 32 torr, PaO₂ 230 torr and bicarbonate 28 mEq/L, showing respiratory alkalosis combined with mild metabolic alkalosis. Continuous high PaCO₂ caused excitability which made assisted ventilation very difficult and PaCO₂ higher and higher. Because of these factors and for the purpose of life-saving, continuous muscle relaxation with pancuronium and sedation with diazepam and morphine were performed, which prevented us from full evaluation of mental state. We tried to decrease intracranial pressure with intravenous furosemide, mannitol and steroid, and used intravenous infusion of sodium thiopental for cerebral protection (Michenfelder and Theye 1973). Throughout the ventilatory insufficiency, we couldn't stop intravenous infusion of dopamine (2-10 μg/kg/min.), suggesting that severe hypercapia resulted in cardiovascular depression. On the 16th day, weaning from ventilator was performed successfully without any problems. Postoperative changes of arterial blood gases and mechanical ventilatory supports were summarized in Table 1. He was completely normal neuropsychologically in the following check at OPD.

DISCUSSION

The prognosis in patients with acute respiratory failure treated with mechanical ventilation has improved since the advent of the respiratory intensive care unit. However, as experience with mechanical ventilation has increased, so has the number of recognized complications. Although some of these are relatively minor, such as nasal necrosis with use of nasotracheal intubation, and mild hyperthermia, others, such as massive gastric distention, tracheal erosion, pneumothorax, and nasocomial pulmonary infection, may be severe and, not rarely, fatal.

The approach to the management of patients with assisted ventilation aimed at the recognition and prevention of complications should lead to a further increase in survival. Such an approach should be based upon knowledge of the spectrum and incidence of complications associated with assisted ventilation, particularly those in which morbidity and mortality have been demonstrated.

Zwilich et al. (1974) has reported the following complications of assisted ventilation in a prospective study of 354 consecutive episodes. The complications attributable to intubation and extubation were prolonged intubation attempt, intubation of right mainstem bronchus, premature extubation and self extubation. The complications associated
with endotracheal or tracheostomy tubes were tube malfunction and nasal necrosis. The complications attributable to operation of the ventilator were machine failure, alarm failure, alarm found off, inadequate nebulization or humidification and over heating of inspired air. The medical complications occurring during assisted ventilation were alveolar hypoventilation and hyperventilation, massive gastric distention, pneumothorax, atelectasis, pneumonia and hypotension.

The complications noticed in this case were pulmonary edema, pleural effusion, pneumothorax and severe acute ventilatory failure (pH 6.90, PaCO₂ 197 torr). Pulmonary edema and pleural effusion probably associated with heart failure were treated successfully with diuretics, fluid restriction, inotropics, colloid, morphine, PEEP and insertion of chest tube. Overpressure applied in airway to overcome aggravating CO₂ retention eventually resulted in pneumothorax. As mentioned previously, severe acute ventilatory failure was mainly caused by airway obstruction resulted from blood clots and debris, secondary to inadequate humidification of inspired air. Inadequate humidification was associated with low dial setting of humidifier, so we could not observe water vapor or droplets in inspiratory rim of ventilator.

The contributing factors to influence the incidence of pneumothorax during continuous ventilatory support are volume-cycled ventilator, chronic obstructive lung disease, excessive tidal volume, increased end-inspiratory pressure, PEEP and percutaneous subclavian venipuncture during pressure breathing (Steier et al. 1971).

Problems with humidifiers during mechanical ventilation may be thermal hazards, electrical and electromagnetic interference problems, contaminants in the humidifier or nebulizer, mechanical or structural failures, inaccuracy in delivered oxygen concentrations, marking or labeling problems, damage caused by the sterilization process, connector failure, human factors, engineering problems, and complaints about the response of manufacturers to reported problems (Bancroft 1982).

Mental changes characteristic of carbon dioxide retention has various correlation to degrees of hypercapnia. In clinical studies of Dulfano and Ishikawa (1965), mental changes occurred at PaCO₂ from 48 to 148 torr, and no obvious correlation could be established between the severity of the mental changes and degree of hypercapnia. In this case, excitability, anxiety and somatic manifestations were observed mental disturbances, but no papilledema and seizure.

The effects of hypercapnia on cardiovascular system may be stimulatory in early and acute stage, but inhibitory in late and chronic stage. The fact that we could stop infusion of dopamine after correction of severe hypercapnia indicates that hypercapnia might cause cardiovascular depression.

The purpose of this report is not to discourage the use of ventilatory support but to alert anesthesiologists who use the technique to the occurrence of complications, the importance of early diagnosis, and means of preventing such complications.

REFERENCES


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수술 후 발생한 환기 부전증

서울대학교 의과대학 마취과학인실 및 홍부의과학인실*

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환로에서 4 증후군이란 전날부터 개심술을 시행한 3세 남자 환자의 수출 후 인공호흡기정에서 심한 급성환기부전증 1례를 경험하였다. 수출후 혈압을 60/40, 녹마상출이 흉부 X-선 소견상 인정되고 인공호흡 시행과정에서 기흉이 발생하였고 급기야는 pH 6.90, PaCO₂ 197 torr의 심한 급성환기부전이 발생하였는 바, 인공호흡기에 부착된 기습기의 조작미숙으로 인한 기습기의 기능저하가 혈액 및 분비물의 응고를 초래시키고 결국은 기도를 폐쇄한 것이 원인이었던 것으로 판단되었으며 기관내 이물질의 장한 음압흡입과 생리식염수를 사용한 세척으로 후유증없이 치료되었다.

본 증례보고의 목적은 인공호흡기 발생가능한 환반중의 종류, 흉부중의 조기진단의 중요성 및 그 예방에 대해 탄탄히에게 다시 한번 경각심을 잃게하는 데 있다.