Negative Pressure Pulmonary Edema: A Case Report

Mohammad Mahbube Mustafa¹*, Kuenza P Wangmo¹, Kamal Ahmed Chowdhury², Sonam Tobgay³, Kuenzang Wangden³, Md. Rafiq ul Alam Talukder⁵

¹Anesthesiologist, Department of Anesthesiology, Jigme Dorji Wangchuk National Referral Hospital, Thimphu, Bhutan
²Junior Consultant, Department of Anesthesiology, Sir Salimullah Medical College & Mitford Hospital, Dhaka, Bangladesh
³ENT Surgeon, Jigme Dorji Wangchuk National Referral Hospital, Thimphu, Bhutan
⁴Resident of Anesthesiology, Jigme Dorji Wangchuk National Referral Hospital, Thimphu, Bhutan
⁵Assistant Professor (Anaesthesia), Sheikh Hasina Medical College, Tangail, Bangladesh

Abstract

Negative pressure pulmonary edema is an uncommon complication of the extubation of the endotracheal tube. An increase in intrathoracic pressure and negative pressure of the lung caused by acute laryngeal spasm results from acute upper respiratory obstruction causing life-threatening pulmonary edema by alveolar-capillary damage is called negative pressure pulmonary edema. We here describe 37-years old male case the preoperative diagnosis of Chronic Bilateral nasal Obstruction associated with headache undergoing Functional endoscopic sinus surgery (FESS) with septoplasty caused negative pressure pulmonary edema while extubation. With the immediate treatment, the patient was discharged without any abnormalities.

Keywords: Extubation, Laryngospasm, Pulmonary Edema

Introduction

Negative Pressure Pulmonary Edema (NPPE) is noncardiogenic pulmonary edema, induce due to an increase in intrathoracic pressure and negative pressure of the lung as a result of acute laryngeal spasm during extubation. NPPE is a life-threatening condition during the extubation after general anesthesia. The prevalence of NPPE occurs in 0.05% to 0.1% [1]. It usually occurs in young healthy patients and during extubation due to which it is also called "post-extubation pulmonary edema" [2]. We here discuss a case of 37 years old male case with a preoperative diagnosis Chronic Bilateral nasal Obstruction associated with headache occurring NPPE during extubation.

Case Presentation

A 37-years old man presented to the Jigme Dorji Wangchuk National Referral Hospital (JDWNRH), Thimphu, Bhutan with complaints of chronic bilateral nasal obstruction associated with headache. Normal past medical history without any comorbidity except, he had hypertension well controlled with medication. There was no history of any past surgery, trauma, allergies, smoking, drug addiction, and infectious diseases. No specific family history was found. On physical examination, there was no abnormality in any system. All the other preoperative assessments including laboratory examinations, imaging examinations, and physical examinations were under normal range.

He was scheduled for Functional Endoscopic Sinus Surgery (FESS) with septoplasty with general anesthesia. The patient was fasted for 12 hours. American Society of Anesthesiologists (ASA) of degree II with height 177 cm, weight 74 kg, BMI 23.6 kg/m² was reported. His Mallampati grade was II degree. Preoperative vital signs were stable with blood pressure of 140/84 mmHg, heart rate 82 bpm, and SpO2 100%.

For Anesthesia induction drugs used Fentanyl 100 μg, Thiopental Sodium 200 mg, midazolam 2 mg and succinylcholine 100 mg then Atracurium 30 mg to maintain muscle relaxant. Operation start time 10:10 am. His vital signs during the operation were stable. EtCO₂ was under normal range. Total operation duration was 2 hours. A total of 1500 mL of ringer lactate solution was administered during the surgery, the urine output was approximately 200 mL, and blood loss was less than 20 mL.

Extubation was done at 12:20 am with self-spontaneous breathing...
observation, tidal volume of 500 L, and when he obeyed the command of opening eyes. Right after extubation patient had breathing difficulties; with mask ventilation positive pressure SpO2 was 88%. But it was ineffective, the further deterioration of SpO2 was observed with 75% to 60%. Pink frothy sputum was found around the mouth and during suction. Atropine 0.5 mg, neostigmine 1 mg was given. Bilateral pulmonary rales were auscultated at the time. With failure to raise in SpO2 patient was reintubated at 12:37 am with sevoflurane 1.8% continuous inhalation, fursemide 20 mg, hydrocortisone 100 mg, aminophylline 0.125 mg, dexamethasone 20 mg, and dexamethasone 8 mg was given. The SpO2 gradually returned to normal and the patient was taken to the HDU with an endotracheal tube. In HDU physical warmer, atracurium 10 mg was used with continuous observation. The blood gas analysis and chest X-ray was taken immediately. The vital signs and urine output was normal in HDU. After 12 hours of observation, the patient was extubated successfully. The patient was sent to the ward without any discomfort and abnormalities.

A chest X-ray is taken when the patient was sent to HDU and after 8 hours. (FIGURE 1 A, B). The chest X-ray at the left side taken when the patient was sent to HDU showed bilateral diffuse, bilateral, hazy, interstitial opacity, pulmonary infiltration throughout both lungs. The chest X-ray at the right side taken after 8 hours patient was sent to HDU showed resolution of the process.

ABG at left side taken when the patient was sent to PACU and right side was taken after 8 hours showed improved SpO2 (TABLE 1).

After the patient was sent to ward ABG report was found under normal ranges, timely vitals, and urine output analysis was done. The blood gas analysis (ABG) and chest X-ray was taken regularly with no suspicion and abnormalities. Then the patient was discharged without any abnormalities after 2 days.

### Table 1. ABG taken after reintubation and after 4 hours

<table>
<thead>
<tr>
<th>Parameters</th>
<th>ABG at PACU</th>
<th>ABG after 4 hours</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base excess (BE)</td>
<td>1.0 mmol/L</td>
<td>1.0 mmol/L</td>
</tr>
<tr>
<td>Calcium (Ca)</td>
<td>1.10 mmol/L</td>
<td>1.10 mmol/L</td>
</tr>
<tr>
<td>Chlorine (Cl)</td>
<td>101 mEq/L</td>
<td>102 mEq/L</td>
</tr>
<tr>
<td>Glucose (Glu)</td>
<td>7.3 mmol/L</td>
<td>7 mmol/L</td>
</tr>
<tr>
<td>Haemoglobin (Hb)</td>
<td>123 g/dl</td>
<td>120 g/dl</td>
</tr>
<tr>
<td>Bicarbonate (HCO3)</td>
<td>24 mEq/L</td>
<td>23.3 mEq/L</td>
</tr>
<tr>
<td>Haematocrit (Hct)</td>
<td>36%</td>
<td>37%</td>
</tr>
<tr>
<td>Potassium (K)</td>
<td>3.86 mEq/L</td>
<td>3.96 mEq/L</td>
</tr>
<tr>
<td>Sodium (Na+)</td>
<td>140 mEq/L</td>
<td>143 mEq/L</td>
</tr>
<tr>
<td>Partial pressure of carbon dioxide (PaCO2)</td>
<td>44 mmHg</td>
<td>42 mmHg</td>
</tr>
<tr>
<td>Acidity (pH)</td>
<td>7.32</td>
<td>7.35</td>
</tr>
<tr>
<td>Partial pressure of oxygen (PaO2)</td>
<td>60 mmHg</td>
<td>100 mmHg</td>
</tr>
<tr>
<td>Oxygen saturation (SpO2)</td>
<td>70%</td>
<td>97.60%</td>
</tr>
</tbody>
</table>

### Discussion

Negative pressure pulmonary edema is an uncommon complication of the extubation of the endotracheal tube. NPPE has been reported mainly by anesthetists as a consequence of postoperative laryngospasm [3]. The pathophysiology of NPPE begins with acute laryngeal spasm leading to severe upper airway obstruction. Intratracheal pressure during acute upper airway obstruction is (> -100 cm H2O) whereas normal inspiratory chest pressure is (-2 cm to -5 cm H2O). The absolute value of intratracheal transpulmonary negative pressure leads to the reduce of alveolar-capillary membrane pressure causing disruption of capillary barrier and alveolar epithelial barrier. All these cause exudation of blood vessels, which makes the red blood cells leak into the alveoli and causes significant lung or bronchial bleeding. These changes lead to red blood cell filtration into the lung and pulmonary edema [4]. The increase in pulmonary blood volume occurs due to the increase in systemic arterial pressure secondary to the release of norepinephrine in response to hypoxia, hypercapnia, and agitation. Frothy pink pulmonary secretions is one of the most important syndrome to diagnose NPPE during extubation and before making the diagnosis of NPPE, other causes of pulmonary edema particularly those requiring a rapid intervention (fluid maldistribution, anaphylaxis, and cardiogenic pulmonary edema), must be considered [5].

The main diagnosis is done by incidence and symptoms, chest radiograph for pulmonary edema, and measure of pulmonary edema fluid/plasma protein ration. With proper treatment, NPPE can rapidly dissipate. The most important treatments include re-establishment of the airway by relief airway obstruction. Adequate oxygenation should be given, encourage the patient to cough after extubation. Application of positive airway pressure via face mask or LMA or Endotracheal intubation with ventilator support. Although NPPE does not result from fluid overload, it is recommended gentle diuresis using low-dose furosemide [6]. The clinical syndromes can usually improve after 12 hours to 24 hours [7]. In our case, the symptoms were improved within 12 hours without any abnormalities. Patients with NPPE history should be kept in mind and prevention methods can be performed accordingly. It is also believed that the incidence of NPPE is higher than that found in the literature due to the fact that the cases are poorly diagnosed and, in many cases, confused with pulmonary edema due to fluid overload, aspiration pneumonitis and bronchospasm [8].

### Conclusions

Functional Endoscopic Sinus Surgery (FESS) with septoplasty caused negative pressure pulmonary edema while extubation. With the immediate treatment, the patient was discharged without any abnormalities.

### REFERENCES

Mustafa, et al.: Negative Pressure Pulmonary Edema: A Case Report
