Acute pulmonary edema due to hypoxia during a difficult intubation in a rheumatoid arthritis patient

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We experienced a rheumatoid arthritis (RA) patient who developed acute pulmonary edema by hypoxia in the absence of spontaneous respiratory effort by sufficient muscle relaxation during anesthesia induction.

A 61-year-old, 136 cm, 48 kg female patient was treated at our hospital for a bilateral total knee arthroplasty due to RA involving the knee joints. During preoperative evaluation of the patient, her chest radiograph showed a suspicious bronchitis at right lower lobe costophrenic angle area (Fig. 1A), whereas the other laboratory tests showed no specific abnormality.

The patient was moved to the operating room to receive bilateral total knee replacement. After normal tidal breathing of 100% oxygen 6 L/min for 3 min using a tightly fitting face mask, anesthesia was induced by injecting 60 mg of propofol and 0.3 μg/kg/min of remifentanil and 30 mg of rocuronium. Under mask ventilation, intubation was performed after 3 min. Intubation was performed using Fiberoptic intubation due to difficulties with endotracheal intubation based on the preoperative c-spine radiography finding of atlantoaxial subluxation. Fiberoptic intubation was tried for 3 min but failed, the subsequent mask ventilation was performed for 3 min, during which the patient's ventilation was not sufficient but maintained, at that time the SpO2, expired tidal volume and ETCO2 was between 90–95%, 200–400 ml and 40–45 mmHg, respectively. But, a few seconds later, mask ventilation was more inadequate, leading to a decrease in the SpO2 up to 70% and cyanosis. Thus, LMA was immediately inserted for adequate ventilation, and the SpO2 were measured as 90%, the patient's ventilation was smooth and SpO2 increased to 100%, whereupon LMA-guided fiberoptic intubation was successfully performed. Five minutes later, a pinkish foamy secretion was observed in the endotracheal tube, and pulmonary edema was shown by a portable chest radiograph; however, no other abnormality was observed (Fig. 1B). Arterial blood gas analysis (ABGA) at an inspired oxygen fraction of 0.5 showed the following: pH 7.3, PCO2 45 mmHg, PO2 149 mmHg, and BE −2.5 mmol/L. Judging that the pulmonary edema was not serious based on the ABGA and chest radiograph, we proceeded with the surgery as planned and intermittent the endotracheal tube suction for 30 min after initiation of the surgery, until the secretion stopped. The total amount of secretion was approximately 20 ml.

The surgery was completed successfully after 4.5 h, and the patient was transferred to the post-anesthesia care unit. Upon improvement of pulmonary edema, verified using chest radiographs, the patient was transferred to the recovery room (Fig. 1C). The patient was moved to the general ward, where she did not show any signs of dyspnea or hypoxia. The chest radiograph taken on postoperative day 2 did not show any abnormality (Fig. 1D), and the patient was discharged from the hospital after 2 weeks.

The type of pulmonary edema frequently associated with airway management or general anesthesia is negative pressure pulmonary edema (NPPE), which is mainly caused by upper airway obstruction such as tracheal extubation, laryngospasm and airway tumor and high negative intrathoracic pressure of −50 to −100 cmH2O generated by respiratory effort, resulting in an abrupt increase in the venous return, leading to increased ventricular volume. This, in turn, increases the left ventricular end-diastolic pressure, leading to pulmonary edema [1]. In the
In the present case, however, no negative intrathoracic pressure was observed during the absence of spontaneous respiratory effort, which was induced by the injection of an intubation dose of a muscle relaxant during the induction of general anesthesia. The patient, thus, was exposed to hypoxia, which resulted from the 6 min insufficient ventilation that was associated with difficulties in intubation until adequate ventilation was achieved through LMA and pulmonary edema occurred immediately after this incident. Therefore, the cause of pulmonary edema in this case is believed to be hypoxia rather than NPPE.

High-altitude pulmonary edema (HAPE) is a type of pulmonary edema associated with hypoxia [2]. In hypoxia-induced HAPE, edema occurs in interstitial and alveolar spaces. HAPE is caused by poor ventilator response, increased sympathetic tone, exaggerated and uneven hypoxic pulmonary vasoconstriction, and inadequate production of hormonal mediators such as nitric oxide [3]. This case report involving pulmonary edema may have been induced, similar to that in the case of HAPE, by the increase in systemic vascular resistance and LV afterload, because of the increased sympathetic tone arising from the continuous stimulation of airway and hypoxic stimulation during intubation failure. Additionally, pulmonary artery constriction, which was triggered by alveolar hypoxia, resulted in an increase in pulmonary vascular resistance, leading to pulmonary hypertension [4]. This change may also indirectly result in an increase in the right ventricular pressure, leading to interventricular septal motion, causing a left ventricular diastolic dysfunction and contributed to the occurrence of pulmonary edema. In addition, a previous study using rats showed that hypoxia inhibits the resorption of fluid in the lungs through the inhibition of Na channels [2]. Thus, similar to that in HAPE, the above-mentioned multiple mechanisms may have caused acute pulmonary edema through hypoxia during anesthesia induction.

RA is primarily a systemic autoimmune disease involving the joints. However, reports have shown that the rate of lung involvement in RA patients is as high as 70% [5]. Thus, RA patients are at risk to changes in the pulmonary conditions such as pulmonary fibrosis, lung nodules, bronchiolitis, tuberculosis, pleural effusions, and bronchiectasis. For these reasons, RA patients are believed to be at high risk for pulmonary injury.

In the present case, acute pulmonary edema improved without administering any specific treatment. The sympathetic tone stimulated by the airway and hypoxia was reduced by the application of inhalation anesthetics and opioids. Further, pulmonary edema improved due to prompt correction of hypoxia, thus, preventing further progression of the condition.
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