Central Neurogenic Hyperventilation Secondary to a Critic Thyroid Status after Aortoaoctic Bypass: A Peculiar Case Report

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Abstract

Central neurogenic hyperventilation (CNH) is a rare condition and defined as a syndrome comprising normal or elevated arterial oxygen tension, decreased arterial carbon dioxide tension, respiratory alkalosis with hyperventilation even during sleep, and the absence of a peripheral respiratory stimulus. The diagnosis of CNH requires the exclusion of pulmonary, cardiac, metabolic-immunological disorders and some medicines that can result in hyperventilation. We detailed the case of CNH in the 4th day after vascular surgery probably secondary to acute metabolic acidosis with hyperlactatemia, due to severe hypothyroidism in the context of critically ill patient.

Keywords
Lactic Acidosis, Metabolic Acidosis, Myxedema Coma, Central Neurogenic Hyperventilation

1. Introduction

Central neurogenic hyperventilation (CNH) was defined for the first time by Plum and Swanson in 1959 as a syndrome with decrease in PaCO₂, increase in PaO₂ and respiratory alkalosis with no other clinical or radiological feature of pulmonary disorder and usually results from lesions in the medial pontine tegmentum and disruption of cortical inhibitory effects of the medullar respiratory center [1].

The main causes of hyperventilation are: pulmonary embolism, severe asthma attack, pneumothorax, myocardial ischemia, congestive heart failure, hyperthyroidism, pheochromocytoma, alcohol and drugs, severe metabolic acidosis, fever, acute pain, ischemia and traumatism. Even if most of CNH are caused by
CNS tumours, especially lymphomas, invasion of brain tissue may produce tissue acidosis and develop hyperventilation [2] [3] [4].

In our literature research we found 8 cases of postsurgical CNH: 5 of them were after the third endoscopic ventriculostomy, 1 after anaplastic astrocytoma resection and 2 after general anesthesia in non-neurosurgical procedures [5]-[12].

2. Case Report

We report a case of a 63-year-old man scheduled for elective aorto-aortic bypass graft surgery due to a 70 mm aortic aneurysm. The patient suffered from hypertension and dyslipidemia and he was in treatment with ARBs and statins. There had been no reports of previous cerebrovascular diseases.

During the procedure, a hemorrhagic shock took place and the patient was transfused with: 8 red blood cell units, 1500cc plasma and 4 grams of fibrinogen. He arrived in the intensive care unit hemodynamically stable and breathing on a venturi mask. The arterial gasometry values at the moment were: pH 7.20, pO2 90, pCO2 45, HCO3 15, BE-7, lactate 8.5, SpO2 96%.

Acidosis was resolved within the next 24 hours. During the early postoperative phase, an acute renal failure took place with creatinine values of 2.8 and uremia of 120. He also suffered a paralytic ileus that was easily solved with IV prokinetics. Lactate levels stayed increased between 2.5 - 3, and hemoglobine was 10 gr/dl.

Four days after surgery, the patient started to feel sleepy, hypothermic and developed an important cognitive impairment. His breathing rate was 50 bpm and he didn’t answer to easy orders. The arterial gasometry values with 4 lpm nasal cannula support were:

pH 7.60, pO2 80 mm Hg; pCO2 16 mm Hg; HCO3 18 mmol/L; BE-10 mmol/L; Lactate 4, SpO2 95%.

A cranial CT-scan and MRI were performed along with an echocardiogram, chest X-ray, D-dimer and biochemical analysis. The results were normal with the exception of: creatinine 1.5, urea 47, decrease in phosphate levels 1.5 - 2.1 mg/DL (normal values 2.5 - 4.5 mg/DL), TSH 0.001 microUI/mL (nv 0.270-4.200) and T4 0.1 ng/dl (nv 0.93 - 1.70).

400 mg IV Hydrocortisone was administered every 24 hours in order to avoid an acute adrenal crisis. In addition, a bolus of 300 microgr IV levothyroxine was given, followed by daily doses of 100 microgr to treat hypothyroidism.

A normal cognitive level, breathing pattern and lactate values were achieved within the next 24 hours.

3. Discussion

The pathophysiologic mechanism of CNH is unclear. It has been shown that in animal models, stimulation of the lateral parabrachial nucleus increased the respiratory rate.

Stimulation of respiratory control areas in thepons and medulla could explain
CNH but it is not sufficient. It has been known that pontine respiratory group neurons modulate the respiratory rhythm. But animal models that disconnect the pontine respiratory group from the medulla have not resulted in CNH.

Hyperlactatemia physiopathology as a cause of CNH is not well known. It is suspected that bicarbonate produced as a result of high lactic acid levels could cross blood brain barrier, resulting in metabolic alkalosis and increasing CO$_2$ levels. As a result, the medullary chemoreceptors responsible for the development of hyperventilation are activated.

The start of a critically ill patient hypothyroidism in just 4 days is not common. Our patient did not need support of mechanical ventilation, benzodiazepines administration or other drug that could affect the thyroid axis. As analgesic drug we used IV fentanyl, this may have decreased the severity of CNH as the publications of Adachi YU et al. have suggested [13], it could be a good palliative treatment in CNH secondary to advanced tumours.

4. Conclusions

The diagnosis of CNH is dependent on the awareness of clinicians to investigate the etiology of tachypnea with or without electrolyte abnormalities. It is important to consider this option in our differential diagnosis when we confront a postsurgical patient after major surgery with poly transfusion and organ failure. Poly transfusion and organic failure with metabolic acidosis after major abdominal surgery is not uncommon, so we must be alert to clinical signs.

As a conclusion, we report Hypothyroidism Coma in critically ill patient showing up as CNH, as an important postoperative complication in abdominal aortic aneurysm surgery. Probably this could be the first CNH case reported in this context at the time.

References


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