Pneumomediastinum in an Accidental Chlorine Gas Exposed Victim

Muhammad Akbar Baig and Maaz Salahuddin

Department of Emergency Medicine, The Aga Khan University Hospital, Karachi, Pakistan

ABSTRACT

Chlorine is an irritant gas, which is implicated in inhalational exposures and can affect the respiratory system leading to complications. We report a case of a 38-year man who presented in the Emergency Department (ED) after exposure to chlorine gas due to an industrial accident. During the course of ED stay, he developed gradual difficulty in breathing, which on investigation, was found to be related to the complication of pneumomediastinum. The patient required endotracheal intubation, but was difficult to ventilate. Bilateral chest tube insertion was performed, which led to the resolution of the pneumothoraces. The emergency physicians should be aware of such a case so that they can intervene.

Key Words: Chlorine, Emergency, Pneumomediastinum, Toxicity.

INTRODUCTION

Chlorine is a yellowish-green irritant gas, which is slightly water-soluble and almost two times heavier than air. Chlorine agents are frequently used for industrial and household chemicals, and for water purification in city water sources.

The respiratory system is the most adversely affected of all organ systems by chlorine gas exposure. Complications can include acute respiratory distress syndrome (ARDS), respiratory failure, pneumomediastinum and death.

In this paper, we report a rare complication of pneumomediastinum in a victim among various patients that presented after accidental chlorine gas exposure caused by gas leakage at a chemical plant in Karachi's industrial zone.

CASE REPORT

A 38-year gentleman presented to the Emergency Department (ED) after an industrial accident of chlorine gas leakage, resulting in hospitalisation of 50 victims. The patient presented with complaints of cough and chest heaviness. Vital signs on arrival were: blood pressures: 138/94 mmHg, heart rate: 120 beats per min, respiratory rate 24 breaths per minute, oxygen saturation 98% on room air. He was afebrile. His systemic examination revealed decreased bilateral breath sounds with poor inspiratory efforts.

There were no significant central nervous system, cardiac or abdominal findings. The patient was nebulised and provided non-invasive mechanical ventilation (NIMV).

His initial set of laboratory tests was sent (Table I). Initial chest radiograph performed was unremarkable and a 12-lead electrocardiogram (ECG) showed sinus tachycardia.

Table I: Initial laboratory investigations.

| Parameters            | Values     |
|-----------------------|------------|
| pH                    | 7.37       |
| pCO2                  | 37.4 mmHg  |
| pO2                   | 103 mmHg   |
| Bicarbonate           | 21 mEq/L   |
| Base Excess           | - 3.6      |
| Oxygen Saturation     | 97 %       |
| Hemoglobin            | 15.4 g/dL  |
| Hematocrit            | 46.7%      |
| White Blood Cell Count| 14.0 × 109/L|
| Neutrophil            | 92%        |
| Lymphocyte            | 5%         |
| Platelets             | 316000/mm3 |
| Procalcitonin         | 0.29 ng/ml |
| Blood Urea Nitrogen   | 12 mg/dL   |
| Creatinine            | 0.9 mg/dL  |
| Sodium                | 139 mEq/L  |
| Pottasium             | 4 mEq/L    |
| Chloride              | 104 mEq/L  |
| Serum Bicarbonate     | 26 mEq/L   |
| Troponin              | <0.006 ng/ml|

The patient was shifted to the observation area. His regular nebulisation did not seem to alleviate symptoms and the patient continued to remain dyspneic. After two hours of ED...
stay, the patient developed neck and chest pain along with respiratory distress. The evaluation revealed subcutaneous emphysema, and swelling with crepitus, which could be felt bilaterally extending from the neck up to the upper thorax. Immediately, a repeat chest radiograph was performed (Figure 1).

Figure 1: Chest radiograph with arrows (red) highlighting linear lucencies along the mediastinal silhouette. Overall findings suggesting subcutaneous emphysema and pneumomediastinum.

Computerised tomography (CT) of the neck and chest with intravenous contrast was performed to find the cause and extent of the damage process. The findings of CT were extensive subcutaneous emphysema of the neck and pneumomediastinum, pulmonary interstitial emphysema, and a linear air density in close approximation to the right posterolateral wall of the trachea at the level of the superior margin of sternum, raising a possibility of tracheal injury.

During ED stay, the patient developed respiratory distress and became diaphoretic. NIMV trial was unsuccessful. Therefore, ED team planned for a definitive airway. Considering a difficult airway, the patient was intubated using a video laryngoscope with 7.5" endotracheal tube. The patient had mucosal erythema and edema near bilateral lateral glossoepiglottic, aryepiglottic and ventricular folds (Figure 2).

The patient was difficult to ventilate due to expanding subcutaneous emphysema and had to be taken on pressure control support of ventilator with a maximum tidal formation of 250 ml. His next set of arterial blood gases (ABGs) showed severe respiratory acidosis with a pH of 6.89 and pCO₂ of > 135 mmHg. The patient was manually bagged and serial ABGs were done which showed interval improvement in the pH to 7.17 and pCO₂ levels to 69.3 mmHg.

Bilateral tube thoracostomy was performed, which allowed resolution of bilateral pneumothoraces and subcutaneous emphysema.

The patient was admitted to Medical Intensive Care Unit (MICU). Cardiothoracic and ear nose and throat (ENT) teams remained on board, advising to manage the tracheal injury conservatively. After 10 days of MICU stay, the patient was successfully extubated and shifted to the special care unit, where he was mobilised; and after removal of chest tubes, the patient was discharged home in a stable condition to be followed in the clinic.

**DISCUSSION**

Chlorine gas inhalation can cause symptoms that can range from mild to moderate but mortality is rare. Minor exposure to chlorine gas may result in irritation of mucous membranes. Major exposure to chlorine gas may result in cough, difficulty in breathing, chest tightness or other pulmonary symptoms. Our case developed violent coughing and shortness of breath after major chlorine gas exposure as a result of an industrial accident.

Hemodynamic parameters can change and result in tachycardia and tachypnea. Our patient exhibited these findings after a few hours of exposure, which made us to think of complications, such as pulmonary embolism, pulmonary oedema, pneumonitis; and as a rare complication, pneumomediastinum. The patient developed subcutaneous emphysema on the neck region sometime after exposure, which was highly suggestive of pneumomediastinum, and was confirmed on interval chest radiographs.

The pathophysiology behind spontaneous pneumomediastinum has already been described. Trigger factors contributing to an increase in the pressure gradient between the alveoli and the interstitium, result in alveolar rupture resulting in dissection of air into the mediastinum. It may happen after a severe strain of cough. This is also the probable mechanism of pneumomediastinum in our patient, which occurred after a violent episode of coughing.

Our patient required endotracheal intubation followed by bilateral tube thoracostomies, which allowed resolution of pneumothoraces. The interval chest radiographs returned to normal.

ED clinicians must be aware of judging such life-threatening and innocuous complications such as pneumomediastinum as per
the clinical scenario. This case presentation draws much-needed attention to a potentially dangerous complication as a result of an industrial accident.

Chlorine is a potent inhalational toxin and can result in disastrous consequences. ED clinicians should be aware when treating such cases; and should attentively follow such cases with high index of suspicion in order to perform timely interventions and prevent complications.

CONFLICT OF INTEREST:
The authors declared no conflict of interest.

PATIENT’S CONSENT:
The consent of the patient was taken prior to the writing of the manuscript.

AUTHORS’ CONTRIBUTION:
Both the authors are responsible for the concept, literature review, drafting and final approval of the manuscript.

REFERENCES

1. Lewis SN. Simple asphyxiants and pulmonary irritants. In: Goldfrank LR, Flomenbaum N, Hoffman RS, Howland MA, Lewin NA, Lewis SN, eds. Goldfrank’s Toxicologic Emergencies. Ed. 11th, New York: McGraw-Hill Medical Pub. Division; 2019:1652-62.

2. Li JH, Wang ZH, Zhu XJ, Deng ZH, Cai CX, Qiu LQ, et al. Health effects from swimming training in chlorinated pools and the corresponding metabolic stress pathways. PLoS One 2015; 10(3):e0119241. doi: 10.1371/journal.pone.0119241.

3. Wang J, Winskog C, Edston E, Walther SM. Inhaled and intravenous corticosteroids both attenuate chlorine gas-induced lung injury in pigs. Acta Anaesthesiol Scand 2005; 49(2):183-90. doi: 10.1111/j.1399-6576.2004.00563.x.

4. Sever M, Mordeniz C, Sever F, Dokur M. Accidental chlorine gas intoxication: Evaluation of 39 patients. J Clin Med Res 2009; 1(5):274-9. doi: 10.4021/jocmr2009.12.1283.

5. White CW, Martin JG. Chlorine gas inhalation: human clinical evidence of toxicity and experience in animal models. Proc Am Thorac Soc 2010; 7(4):257-63. doi: 10.1513/pats.201001-008SM.

6. Morim A, Guldner GT. Chlorine gas toxicity. 2020 Jun 30. In: Stat pearls. Treasure Island (FL): Stat pearls publishing; 2020 Jan.