Chemical Pneumonia Due to Paint Thinner Ingestion: A Case Report and Literature Review

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Abstract

Paint thinner is an organic solvent which includes aromatic hydrocarbons and is widely used in the paint, varnish, and plastic products industries. Regular inhalation of such solvents is known to cause chronic intoxication. Oral thinner over-exposure is rare but quite fatal. Volatile liquids can cause pulmonary complications even with oral ingestion. Therefore, in patients with oral volatile liquid thinner intake, emergency physicians should be aware of local complications as well as the systematic effect even with initial normal physical or radiological findings. In this report, we present a patient with chemical pneumonia due to self-vomiting after accidentally drinking paint thinner.

Keywords: Chemical pneumonitis, thinner, hydrocarbons, ingestion

Introduction

Exposure to hydrocarbons is common in modern society. Thinner is an organic solvent that contains aromatic hydrocarbons such as toluene and xylene and is easily accessible in products such as gasoline, paint, varnish, and plastic production industries. Types of exposure include unintentional ingestion, intentional recreational abuse, unintentional inhalation, and dermal exposure or oral ingestion in a suicide attempt. Regular inhalation of such solvents is known to cause chronic intoxication. Toluene and xylene are absorbed in the gastrointestinal tract and are shown to cause rhabdomyolysis, metabolic acidosis, hepatic and renal dysfunction, neurotoxicity, and pulmonary toxicity due to their lipophilic affinity in rodents and humans.

Oral thinner over-exposure is rare, but quite fatal. Oral intake of 45-50 mL thinner is known to cause severe complications (1). Toluene accumulates in the liver following oral exposure, and thinner accumulates in the brain following respiratory exposure. Volatile organic compounds act as central nervous system depressors due to their lipophilic characteristics and cause death after oral ingestion. Herein we report an adult male who was admitted to our hospital with complaint of fever with the history of paint thinner ingestion.

Case Report

A 54-year-old male patient presented to the emergency department (ED) with fever. He had drunk half bottle of paint thinner (approximately 100 mL) accidentally 9 hours before his admission. Upon realizing his mistake, he had self-induced vomiting in an attempt to get rid of the thinner about 2-3 times, and then had drunk a lot of water to minimize the effect of the thinner.

Physical examination revealed an alert and oriented male with a regular pulse of 110 beats/min, blood pressure of 119/75 mmHg, respiratory rate of 20 breaths/min, body temperature of 38.9 °C, and O₂ saturation of 98% on room air. Cardiac examination revealed normal S1 and S2 with no audible murmurs, rubs, or gallops. Chest examination showed bilateral equal aeration; there were crepitant rales over the right lower zone but no wheezing or rhonchi. The abdomen was soft, non-tender, and non-distended, without any evidence of organomegaly. No lower
Extremity edema, cyanosis, or clubbing was seen. The peripheral arterial pulses were palpable. The neurologic examination was normal.

Laboratory tests, including arterial blood gas, renal and liver function tests, and a comprehensive metabolic panel were all within normal limits. The results of the complete blood count were as follows; white blood cell count: 19.300 L; neutrophil ratio: 83.4%; platelet count, hemoglobin and hematocrit levels were within the normal range. No invasive or infiltrative lesion was detected in the chest X-ray (Figure 1).

The patient was hospitalized for supportive treatment and further workup and monitoring. At third hour of admission, the patient developed an attack of bronchospasm. Meanwhile, the vital findings were as follows: SpO₂: 97% (on room air), pulse: 109/min, blood pressure: 120/67 mmHg. After inhalation of nebulized ipratropium bromide plus budesonide, his symptoms regressed. At 7th hour of admission, the same inhaler treatment was administered because of a recurrent attack of bronchospasm. Therefore, chest computed tomography (CT) scan (without contrast) was performed for differential diagnosis. It showed consolidation areas containing air bronchograms secondary to parenchymal infiltration in the inferior lobe of the right lung (Figure 2). Hence, chemical pneumonitis was diagnosed and amoxicillin-clavulanate was started. Patient was discharged after he had been symptom-free and hemodynamically stable for 24 hours on the 3rd day of admission.

Discussion

Oral intake or inhalation of less than 1 mL of some hydrocarbons has been shown to cause chemical pneumonia and death. Respiratory symptoms generally begin in the first few hours after exposure and usually resolve in 2-8 days. Symptoms such as cough and broncho-obstruction may occur shortly after oral intake. Tachypnea, wheezing, and chemical pneumonitis may follow thereafter. In such cases death due to chemical exposure, which is usually related to bacterial infections and other respiratory complications, may ensue (2). Our patient, who had self-vomited after accidentally having drunk thinner, developed fever and respiratory signs. Inducing vomiting in such patients is not advised because it can increase the risk of pulmonary complications as a result of aspiration (3).

Hydrocarbons can be aspirated after oral ingestion and may cause toxic effects on lungs. Nonetheless, they may also cause systematic toxic effects due to oral, respiratory, or dermal exposure. Especially in thinner and naphtha intoxications respiratory symptoms can slowly occur. Death has been reported in patients with multi-organ failure secondary to oral intake of hydrocarbons (4). While signs of chemical pneumonitis can appear in the chest radiogram as early as 30 minutes after exposure to any chemical in symptomatic patients, they may variably occur between 2-24 hours in asymptomatic patients (5). In our case, fever and leukocytosis with negative chest X-ray occurred 9 hours after the oral intake and signs of chemical pneumonitis in the chest CT settled 16 hours after the intake. Therefore, the patient was given broad-spectrum antibiotic due to increasing infiltrate in radiological imaging. Occurrence and persistence of fever, increasing infiltrate in chest radiograph, leukocytosis or sputum or tracheal aspirate positive for bacteria after hydrocarbon aspiration over 48 hours suggests bacterial superinfection. Hence, pneumonitis caused by hydrocarbon aspiration should not be treated routinely with antibiotics unless signs of secondary infection (6).

In the literature reports fatal toluene levels were reported to be 29-119 µg/g in 3 cases (7); in one case who was externalized in...
full health after 8 days of follow-up in the intensive care unit, toluene level was reported to be 17 µg/g (1). Blood toluene level test cannot be performed in our hospital therefore, we cannot provide any data about the blood toluene level of our case.

In a report of 20 cases with toluene inhalation, the most frequent complaint was muscle weakness due to hypokalemia, followed by altered mental status and gastrointestinal complaints including nausea, vomiting, and abdominal pain with elevation of creatinephosphokinase concentration to the level approximately 6 times the upper normal limit and elevated gamma-glutamyl transpeptidase and alkaline phosphatase (ALP) levels (8). In another report of 37 adult thinner intoxication cases, the most frequent complaint was nausea and vomiting with elevated ALP and lactate dehydrogenase levels, approximately 2-3 times the upper limit of normal (9). In our case, there was no abnormality in the electrolytes, or liver, or kidney function tests.

Treatment with surfactant in pediatric patients with acute respiratory distress syndrome (ARDS) due to toluene inhalation and treatment with budesonide and nitric oxide in patients with respiratory distress and bronchospasm were found effective (5). On the contrary, treatment with corticosteroids in dogs with fulminant hydrocarbon aspiration was found ineffective (10). Our patient had fever without respiratory distress in initial examination and had bouts of bronchospasm 12 hours and 16 hours after exposure. The patient was treated with inhalation of nebulized ipratropium bromide plus budesonide during the bronchospasm attacks.

Emergency physicians should be aware of pulmonary complications even with oral exposure to volatile liquids. Therefore, the occurrence of local pulmonary complications as well as the possible systemic complications should be closely monitored due to implication in pneumonitis but also in central nervous and gastrointestinal system toxicity, arrhythmias, hypokalemia and metabolic acidosis. The absence of symptoms or radiological findings immediately after thinner or any chemical ingestion or the presence of only short-lived symptoms is possibly determined by the amount and physical characteristics of the substance reaching tracheobronchial tree. Herewith, the healthcare professionals should be aware of developing serious complications in patients with chemical exposure who have normal physical findings initially. It should not be forgotten, in 5% of cases with pneumonitis, the disease progresses rapidly with severe manifestations such as multiple organ failure and ARDS (11).

Conclusion

The patients with a history of any chemical exposure should be hospitalized for at least twenty-four hours after exposure for monitoring of local and systemic toxic effects. It should be noted that the diagnosis of hydrocarbon exposure is based upon clinical features. It is not necessary to take the results of the laboratory tests rapidly because they do not change the management priorities.

Ethics

Informed Consent: It was taken.

Peer-review: Externally peer-reviewed.

Authorship Contributions


Conflict of Interest: No conflict of interest was declared by the authors.

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References