Massive Myocardial Infarction Due to Acute Left Main Coronary Artery Occlusion in a Young Organophosphorus Poisoning: Case Report

Gautam Archana¹, Patil Vijaysinh², Rayannavar Nikhilesh³, Goel Himalaya⁴, Bholegare Sakshi⁵, Bammidi Rohit⁶

¹Head of the Department of Critical Care Unit, Krishna Institute of Medical Sciences, Karad, Maharashtra, India; ²Interventional Cardiologist, Krishna Institute of Medical Sciences, Karad, Maharashtra, India; ³,⁴,⁵,⁶Junior residents, Department of Internal Medicine, Krishna Institute of Medical Sciences, Karad, Maharashtra, India.

ABSTRACT

Introduction: In the farmlands what has been found to be a common chemical for suicidal attempts are Organophosphate poison. This is common in many of the countries. Organophosphate poison is quite dangerous and can have impact once someone consume. Even the smell of this liquid or touch of the chemical to the skin can have a very harmful effect. Fatal left main carotid artery occlusion is rarely reported in literature.

Aim: To study the case of consumption of Chlorpyriphos- cypermethrin combination

Case Report: In the present paper we have reported a specific case of a person whose age was 25 years and who consumed a Chlorpyriphos- cypermethrin combination who later developed massive myocardial infarction with an ST-elevation >12 mm in aVR followed by a cardiac arrest.

Discussion: Cardiac abnormalities are rare in organophosphorus compound poisoning. The patients have include arrhythmia, heart failure, cardiogenic shock, and sudden death. In those patients, clinical manifestations of damage caused by organophosphorus pesticides to the myocardium vary.

Conclusion: During the management of organophosphorus compound poisoning, intensive cardiac monitoring should be considered.

Key Words: Massive Myocardial Infarction, Acute LMCA Occlusion, Organophosphorous, Poisoning, Cardiac, Pyrethroid compound, Cypermethrin

INTRODUCTION

There are many types of inhibitors available, one of them are Organophosphate, it is potent cholinesterase inhibitors, which is capable of causing severe cholinergic toxicity following cutaneous exposure, inhalation or ingestion.¹ Chlorpyriphos is a pesticide that contains a broad-spectrum chlorinated organophosphate.² We use that to control many kinds of pests, termites, mosquitoes and roundworms. According to world health organization Chlorpyriphos is considered moderately hazardous on the base of acute toxicity. Exposure surpassing recommended levels has been linked to neurological effects, persistent developmental disorders, and autoimmune disorders. The symptoms of severe poisoning is seizures, unconsciousness, paralysis and suffocation from lung failure.³ A pyrethroid compound Cypermethrin widely used as an insecticide, is moderately toxic but excessive exposure can cause nausea, headache, muscle weakness, salivation, shortness of breath and seizures. Apart from the drug toxicity, these compounds cause many cardiac complications like QT prolongation, conduction blocks, arrhythmia. But rarely do they cause myocardial infarction.

CASE REPORT

A male who is 25 years old was admitted to the intensive care unit with of alleged history of intentional consumption of HAMLA (Chlorpyriphos 50% + Cypermethrin5%) around 50 ml followed by 3 episodes of vomiting. The patient was
Archana et al: Massive myocardial infarction due to acute left main coronary artery occlusion in a young organophosphorus poisoning

brought to the hospital within 3-4 hours of consumption. On examination he was drowsy, sweating profusely with GCS of 6 (E1 V2 M3), pupils were constricted <2mm and neck holding was absent. Peradeniya organophosphate score was 7, suggestive of moderate poisoning. Vitals at the time of admission were was-PR-60/min; BP-110/70mmHg, RR-32/min; SPO2- 92%. Patient had no comorbid conditions. The patient was immediately taken on mechanical ventilation (ventriculocoronary arterial communication) VCAC mode and was treated with atropine, Pralidoxime and other supportive measures (IV fluids, gastric lavage, savlon body wash, and activated charcoal). Arterial blood gas analysis suggested metabolic acidosis with compensatory respiratory alkalosis and the blood analysis revealed a serum cholinesterase level of 1858 IU/L with serum CPK NAC level of 229 IU/L. ECG at the time of admission had sinus rhythm with a heart rate of ~ 68/min. For the next 3 days, the patient was kept on mechanical ventilation under mild sedation with RASS of -2 (Richmond Agitation-Sedation Scale) and heart rate was maintained at ~100/min on atropine at 0.2mg/hr. Inj. Heparin 5000 IU I.V 12 hourly was started as DVT prophylaxis on day 3. On day 4 around 10.30 P.M, the patient developed sweating and sinus tachycardia with a heart rate of 140/min lasting for 3-4 hours. Sudden ECG changes were noticed in the cardiac monitor and the ECG recorded suggested acute left main coronary artery occlusion with deep ST depression >10 mm in anterior and inferior leads and ST-elevation around 12 mm in aVR. Sooner he developed cardiac arrest and could not be revived by CPR and inotropic supports. Biochemical cardiac markers sent at that time were positive with Serum Trop I of 3.5 ng/ml.

Table 1: Laboratory parameters

<table>
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<th>Day</th>
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<tbody>
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<td>Hb (g/dl)</td>
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<td>13.7</td>
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<tr>
<td>Total count(/µl)</td>
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<td>Creatinine(mg/dl)</td>
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<td>142</td>
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<td>Potassium (mEq/l)</td>
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<td>Serum troponin (ng/ml)</td>
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<td>1155</td>
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<td>CPK NAC</td>
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<td>1481</td>
<td>&gt;1000</td>
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Table 1: (Continued)

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<td>Lactate</td>
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<td>0.7</td>
<td>0.5</td>
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Urine routine, LFT Normal

Figure 1: ECG on admission. Sinus rhythm with heart rate ~68/min.

Figure 2: ECG on day 4. Sinus tachycardia with heart rate ~140/min ST elevation in aVR >12mm, ST depression in I, II, III, aVF, V₂ to V₆ >10 mm.

DISCUSSION

Cardiac abnormalities are rare in organophosphorus compound poisoning. The patients have to include arrhythmia, heart failure, cardiogenic shock, and sudden death. In those patients, clinical manifestations of damage caused by organophosphorus pesticides to the myocardium vary. Myocardial injuries and myocardial infection are rarely reported, which cause by organophosphorus poisoning.

According to a study conducted by Aktar S et al. in 2019 on the complications of Organophosphorus poisoning, out of 100 cases, 29 had cardiac complications like premature ventricular contractions, prolonged QT, conduction block,
polymorphic VT, Torsade de Pointes, sinus bradycardia which occurred during early hours of acute poisoning. But none of them had myocardial infarction as a complication in their study. For the development of these complications Hypoxia, electrolyte imbalance and acidosis is very important. Our case did not have any predisposing factors and never developed these complications. (Though metabolic acidosis was present when the admission is going on, it was corrected immediately).

According to a report of a case by P Joshi et al. on a 40-year-old patient with parathion consumption, it was found that patient had sinus bradycardia on admission. On the second day, the patient developed ST-segment elevation with T inversion in V3-V6 with Trop I of 3ng/ml. The patient was treated with antiplatelet, nitrates, LMWH and was discharged with the resolution of ST and T changes. In contrast to this, our case was a young male with no previous comorbidities who developed Massive myocardial infarction on day 4 with ECG showing ST elevation > 12 mm in aVR and ST depression in other leads >10mm, with serum Trop I of 3.5 ng/ml indicating blockade in LMCA territory. Later he had a cardiac arrest and could not be revived.

The mechanism of cardiotoxicity by these compounds is not completely understood.

There are few postulations by which OP compounds can cause cardiac toxicity. 4,5

1. Brief period of increased sympathetic tone.
2. Prolonged period of parasympathetic activity
3. QT prolongation followed by torsades de pointes. And also, other mechanism includes sympathetic/parasympathetic overactivity, hypoxia, dyselectrolemia and direct toxicity. Coronary vasospasm can also lead to myocardial infarction. Coronary vasospasm was caused by intracoronary injection of acetylcholine in individuals as demonstrated in a study conducted by Hario et al. and Yasue et al. 6,7

In a person with previous coronary artery plaque, the release of vasoactive amines like histamines and catecholamines leads to penetration of collagen matrix of plaque-causing erosion and rupture causing myocardial infarction. 8 Release of inflammatory cytokines like histamines, neutral proteases, platelet-activating factors, arachidonic acid products, chemokines causes vasospastic angina, nonvasospastic angina, myocardial infarction known as Kounis phenomenon. 9

Serious cardiac complications cause by Patchy myocardial involvement as a result of direct cardiac toxicity. 10,11 Myocardial may not be manifest clinically or on echocardiography because it is involved in patchy. Continuous cardiac monitoring should be undertaken to detect dynamic cardiac changes. Continuous cardiac monitoring should be undertaken to detect dynamic cardiac changes.

CONCLUSION

Organophosphorus poisoning can cause severe cardiac complications within few hours of exposure. Myocardial infarction occurs as a complication during the early phases of poisoning, but it also has chances of occurring during the recovery phases. Thus, during the management of organophosphorus compound poisoning, intensive cardiac monitoring should be considered.

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REFERENCES

