Fatal Iron Toxicity in an Adult: Clinical Profile and Review

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Abstract

Acute iron toxicity is usually seen in children with accidental ingestion of iron-containing syrups. However, the literature on acute iron toxicity with suicidal intent in adults is scant. We report an instance wherein an adult committed suicide by ingestion of multiple iron tablets. Delay in treatment was there due to misdiagnosis of the intoxicating agent. She developed fulminant hepatic failure with rapid clinical deterioration. Despite aggressive supportive management, the patient succumbed to the toxic doses of iron. Clinical course and postmortem features are discussed with a review of the literature.

Keywords: Ferrous sulfate, hepatic failure, Perl’s stain, toxidrome

Introduction

Iron is the most abundant trace element in the body and is essential in most biological systems.[1] Iron is an intracellular poison, and it exerts its toxic effect on mitochondria by shunting electrons away from the electron transport chain, uncoupling the oxidative phosphorylation, thus leading to anaerobic metabolism.[2] Iron poisoning in an adult is uncommon. Most of the cases of iron poisoning reported in literature are of pediatric age group,[2‑4] while only four cases[1,5,6] were of older age group.

In this article, we present a case of acute iron intoxication in a 20-year-old female who was misdiagnosed as aluminum phosphide poisoning and subsequently managed as iron poisoning. Clinical symptoms, laboratory tests, institutional toxicology screening, and histopathology which corresponded with severe iron intoxication are discussed.

Case Report

A 20-year-old female allegedly consumed some tablets with suicidal intention. She initially presented at primary health center with abdominal pain, vomiting, diarrhea, and shock. As the intoxicating agent was not clear, treatment was started for aluminum phosphide poisoning as it is quite prevalent in the area. Initial complete blood count and serum electrolytes were unremarkable. The patient’s condition deteriorated and she was referred to our hospital on the 3rd day. At our hospital, she was further investigated; hemoglobin was 9.1 g/dl and total white blood cell count was 11,900/cumm. Liver function tests showed marked elevation in the liver enzymes (aspartate aminotransferase [AST] – 404 IU/L and alanine aminotransferase [ALT] – 1982 IU/L). Other investigations showed total bilirubin – 6.37 mg/dL, direct bilirubin – 4.60 mg/dL, indirect bilirubin – 1.77 mg/dL, total protein – 6.43 g/dL, serum albumin – 3.35 g/dL, alkaline phosphatase – 112 IU/L, and amylase – 465 U/L. Prothrombin time was 56 with international normalized ratio of 5.7. Silver nitrate test was negative for phosphide. Her detailed history was sought again, in which she revealed consumption of many ferrous sulfate tablets. Hematocrit profile showed serum iron levels as 185 µg/dl, serum ferritin – 692 ng/dl, total iron binding capacity – 90 µg/dl, and percentage saturation – 2%. Blood gas analysis showed severe metabolic acidosis. Chelation therapy with desferrioxamine was started, and supportive measures were continued. However, her clinical condition deteriorated further, and she died within 48 h of admission due to multiorgan failure.

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Autopsy revealed enlarged liver. Histopathology showed periportal necrosis and bile cholestasis [Figure 1a and b]. Other organs were unremarkable on routine hematoxylin and eosin stain. On Perl’s stain, iron deposits were noted in the liver, lungs, and spleen, suggestive of iron overload [Figures 1c, d and 2]. Cause of death was attributed to multiorgan failure due to iron poisoning.

**DISCUSSION**

The present case discusses delayed clinical presentation of iron toxicity. Iron toxicity has five clinical stages. These stages of clinical course may not be found in all patients, and in cases of massive overdose, the patient may present in shock.[7] Rapidly progressive clinical course from Stage I to Stage III was also observed in reported cases,[4] and this may cause potential misdiagnosis when history is not clear.

Unreliability is inherent in overdose or poisoning histories as was in the present case. In the present case, initial history was misleading and the patient was treated for aluminum phosphide poisoning, as it is quite prevalent in this part of the country,[8] and has similar toxidrome as that of acute iron toxicity. Toxicology screening for phosphide was negative, leading to re-interview of the patient which revealed consumption of multiple iron tablets. Although misdiagnosis of the intoxicating agent was corrected, delay in the treatment of iron toxicity leads to adverse clinical outcome. Patients are reported to have better outcome and survival if they present early and are managed with desferrioxamine, bowel irrigation, and other supportive treatment.

In the present case, it was not clear as to how many iron tablets patient had consumed, and hence, calculation of total iron ingested was not possible. If the patient presents early, then serum iron concentration of 300 mcg/dL or above can present as moderate toxicity, while serum iron concentration of 500 mcg/dL or above represents severe toxicity.[6] Serum iron levels are best estimated within 2–6 h[9,10] as rapid distribution of iron from circulation to tissues results in stabilization of iron levels.[9] Hence, diagnosis was not confirmed by serum iron levels where presentation was after 36 h[9] and 48 h.[1] Other parameters which showed positive correlation with elevated serum iron and hence severe toxicity include severe vomiting or diarrhea, shock, coma, coagulopathy, metabolic acidosis, hyperglycemia, and leukocytosis.[2] Our patient had iron levels of 185 IU/L after 48 h of ingestion, raised AST/ALT levels, deranged liver function tests, coagulopathy, and state of shock indicating severe toxicity.

Hepatotoxicity is known sequela of iron poisoning and is associated with 50% of mortality.[5] It is found to be dose dependent[11] and hence useful in cases which present late, as serum iron levels may not be a reliable prognostic indicator in such cases. In the present case, hepatic toxicity was manifested by periportal necrosis and cholestasis. Coagulative degeneration, shrinkage of hepatocytes, intense cytoplasmic eosinophilia, and karyorrhexis are also reported.[4] In the present case, Perl’s-stained iron deposits were found in hepatocytes. However, pigment did not attribute to “alcoholic iron” as liver was noncirrhotic and deceased was nonalcoholic.[12]

**CONCLUSION**

As suicidal iron tablets overdose is uncommon, it can be misdiagnosed with toxicity by the agent having a similar clinical outcome. However, toxicological assistance to an intensivist is vital in such cases as it rules out concomitant intoxication or intoxication by an agent having similar toxidrome and fatality due to delay in treatment of iron toxicity can be prevented.
Declaration of patient consent
The authors certify that they have obtained all appropriate patient consent forms. In the form the patient(s) has/have given his/her/their consent for his/her/their images and other clinical information to be reported in the journal. The patients understand that their names and initials will not be published and due efforts will be made to conceal their identity, but anonymity cannot be guaranteed.

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Conflicts of interest
There are no conflicts of interest.

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