### A fatal case of multi-organ failure due to acute yellow phosphorus poisoning

#### Abstract

Yellow Pphosphorus is a non-metallic irritant used in various sectors industries such as the rodenticide, firecracker, match, and fertilizer industries. PYellow phosphorus poisoning is responsible for deaths amongfatal in both children and adults. Accidental yellow phosphorus poisoning is frequently reported in children, whereas suicidal consumption is not uncommon among adults. Herein, we present the case of a 30-year-old female patientwoman who ingested Ratol paste containing yellow phosphorus in an attempt to commit suicide. Her initial chief complaints were nausea, vomiting, and diarrhoea during hospitalization hospitalisation, followed by a symptomless phase with stable vitals vital parameters on the second day of hospitalisation. S; she was managed conservatively and discharged. She took discharge against medical advice. She was readmitted to the same hospital after two days on the fourth day after Ratol ingestion with complaintsing of generalized generalised weakness, body pain, drowsiness, loss of appetite, and breathing difficulties. She developed severe complications due totowing to the intoxication and died. An autopsy was performed, and h. Histopathological and the toxicological examinations were carried out. We foundrevealed characteristic features of yellow phosphorus toxicity in her organs. We concluded that the cause of death was hepatic encephalopathy and multi-organ dysfunction syndrome caused by the yellow phosphorus poisoning.

**Keywords**: Phosphorus, Hepatic encephalopathy, Multi<del>ple</del>-organ failure

### Introduction

In India, around approximately 70% of rural households still depend primarily on agriculture for their livelihood [1]. Fre-An unchecked growth of rodents like-such as rats around houses and sometimes in farms and fields can be problematic because  $\frac{\text{they-} rodents}{\text{can}}$  can potentially spread diseases  $\frac{\text{ses}}{\text{that}}$ would hamper the food supply and consequently lead to economic loss. Hence, rodenticides are widely marketed in India, and various preparations are available, such as yellow phosphorus in the form of Ratol paste and powder. This preparation is cheap and easily-readily available in the open market and on online e-commerce sites in India;, this which contributes to the frequently reported cases of suicides due to Ratol ingestion. Accidental The accidental consumption of Ratol paste at home is not uncommon among children. Rodenticides containing 3-5% yellow phosphorus are currently available; and Ratol paste contains 3% yellow phosphorus [2]. It is highly toxic and does not have <u>any an</u> antidote. Accidental poisoning with Ratol paste is very common since it is very similar to toothpaste [3–5]. 3-5 Yellow phosphorus is also used in fireworks and matches, which leads to chronic poisoning in workers [6]. Yellow phosphorus is a kind of non-metallic protoplasmic poison. It is rapidly absorbed from the digestive tract and is primarily metabolized metabolised by the liver 71. The smallest fatal dose is 8 mg; but however, the usual fatal dose is 1 mg/kg 81. The v¥omitus after phosphorus ingestion is luminescent and has a characteristic garlic odour. If the patient survives the initial gastrointestinal irritation phase, hepatic toxicity ensues secondary to systemic poisoning. Herein, we report a fatal case of suicidal yellow phosphorus poisoning.

# **CASE REPORT**

A 30-year-old female patientwoman was admitted to the Department of Trauma & and Emergency Medicine in a tertiary care hospital with complaintsing of a burning sensation in the mouth, nausea, and vomiting. The vomitus had a garlic odour. Her husband stated that she had ingested 10-12 g

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Ratol paste about approximately 2 hours before. Gastric lavage was done with 1:5000 KMnO<sub>4</sub> and activated charcoal was performed. The patient was conscious with a blood pressure of 140/80 mmHg, pulse rate of 84 beats per minute, and respiratory rate of 20 breaths per minute. Results of different tests performed at the time of admission are shown in Table 1.

Bedside <u>abdominal</u> ultrasonography (<u>USG</u>) of the <u>abdomen</u>-was unremarkable. Symptomatic management was <u>done achieved</u> with N-acetylcysteine (NAC). On the second day <u>of admission</u>, ther <u>patient's</u> clinical status improved, and she was asymptomatic. The patient, <u>and-supported by</u> her relative, took-was <u>discharged against medical advicedischarge despite insistent medical advice to remain hospitalized</u>. On the fourth day <u>after Ratol ingestion</u>, the patient returned to the hospital with <u>generalized generalised</u> weakness, body pain, drowsiness, and breathing difficulty. She was <u>drowsypresented</u> with hypotension (blood pressure of 88/58 mmHg), tachycardia (110 b<u>eats per minutepm</u>), and tachypnoea.

On general examination, icterus was present. Laboratory workup results fromen the fourth day are shown in Table 1. A bedsideBedside abdominal ultrasonographyUSG showed hepatomegaly and fatty changes. Hence, a provisional diagnosis of yellow phosphorus poisoning with hepatic encephalopathy and multi-organ failure was made. On the fifth day, the patient developed suddenonset bradycardia and hypotension, which rapidly led-progressed to the patient's death with no opportunity to perform liver transplantation. An autopsy was conducted performed.

### **AUTOPSY FINDINGS**

On postmortempost-mortem examination, scleral icterus was present. Both pleural cavities were filled with approximately 350 ml-ml of haemorrhagic fluid each. Interlobular fissures of both lungs showed petechial haemorrhages. The right and left lungs weighed 410 g and 455 g, respectively (RV: 450 g and 375 g, respectively). The cut section showed that the lungs were congested. Pinpoint haemorrhages were present over the heart surface. The weight of the heart weighed was about approximately 270 g (RV: 243 g). The peritoneal cavity was filled with 750 ml-ml of haemorrhagic effusion. The stomach mucosa was haemorrhagic. Pinpoint to pinhead-sized petechial haemorrhages were present over the mesenteries, liver, and kidneys (Figure 1A, 1B). Yellowish A yellowish discoloration of thewas present on the liver, kidneys, and brain surfaces was present. The weight of the liver was about 1100 g (RV: 1100 g); the weights of the right and left kidneys was was were 114 g and 129 g, respectively (RV: for both 288 g); and the weight of the brain was 1177 g (RV: 1233 g). The cutCut sections of the liver (Figure 1C) and kidneys (Figure 1D) showed pinpoint haemorrhages.

Figure 1. Gross examination of the: (A)— liver showing pinhead-sized haemorrhages over the surface and yellowish discoloration; (B)— kidneys showing pinpoint haemorrhages over the surface and yellowish discoloration; (C)— cut surface of the liver depicting pinpoint haemorrhages within the hepatic parenchyma; (D)— cut section of the kidney depicting pinpoint haemorrhages and yellowish discoloration of the renal parenchyma.

Histopathological examination of the lung-showed acute interstitial inflammatory cell infiltrates along with multiple focal alveolar haemorrhages in both lungs (Figure 2A, 2B). The heart showed focal necrotic fibres with focal acute myocarditis (Figure 2C).

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- •All Figures, Schemes and Tables should be inserted into the main text close to their first citation and must be numbered following their number of appearance (Figure 1, Scheme I, Figure 2, Scheme II, Table 1, etc.).
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### Figure 2.

Photomicrographs of the (<u>A and B</u>) lung (<del>A and B)</del> and (<u>C) the</u>-heart (<del>C)</del>. (<u>A</u> and <u>B</u>)—Inflammatory cell infiltrates along withand focal alveolar haemorrhage (H&E, 4X and 10X, respectively); (<u>C</u>)—Focal necrotic fibres with neutrophilic infiltrates, suggestive of focal acute myocarditis- (H&E, 10X).

The liver showed non-zonal necrosis, karyorrhexis, vacuolization intracellular bile pigment deposition, mild periportal inflammation, and piecemeal necrosis (Figure 3A, 3B). The kidneys showed vacuolization vacuolisation of proximal tubular cells and multifocal necrosis of the lining epithelial cell linings with sparing of the glomeruli (Figure 3C). The pancreas showed focal fat necrosis along with necrosis of large areas of the pancreatic parenchyma (Figure 3D). Histological analysis of the bone marrow was not carried outperformed. All features were suggestive of multiorgan failure comprising submassive hepatic necrosis, focal acute myocarditis, acute renal tubular necrosis, and pancreatic necrosis typical of phosphorus poisoning, which was confirmed after a qualitative chemical analysis report showed the presence of phosphorus in tissues sampled during the autopsy.

## Figure 3.

Photomicrographs of the (A, B) liver (A, B), (C) kidney (C), and (D) pancreas (D). (A)— Non-zonal necrosis, karyorrhexis, vacuolizationvacuolisation, and piecemeal necrosis (H&E, 4X);, [B]— Vacuolization in liver tissue (H&E, 10X); (C)— Vacuolization Vacuolisation of the proximal tubular cells and multifocal necrosis of the lining epithelial cells with sparing of the glomeruli (H&E, 10X); (D)— Parenchymal necrosis of the pancreas (H&E, 10X).

## DISCUSSION

Yellow phosphorus is a toxic substance that is used in matches, fireworks, and rodenticides [9]. Developing Several cases of intoxication with yellow phosphorus in developing and underdeveloped countries have been reported intoxication with yellow phosphorus; but however, this is rarely reported in developed countries. In developing countries, the intoxication generally results from accidental oral ingestion, although suicidal ingestion is also not uncommon. 10,11

Ratol paste contains 3% yellow phosphorus, a far more toxic substance than red phosphorus [12,13]. 12,13 Yellow phosphorus is categorized\_categorised as a highly lethal rodenticide when ingested in doses exceeding 1 mg/kg. In our case, the deceased ingested approximately 10-12 g of yellow phosphorus rodenticide. As reported in the literature, victims of Patients with yellow phosphorus poisoning may be initially symptomatic; however, recovery is observed after 2-3 days of ingestion., but later Nevertheless, symptoms of acute liver failure subsequently develop [12]. 12 In our case, the patient had a similar progression of complications. Hence, we surmise that patients with acute yellow phosphorus poisoning should be monitored closely for a 1 week since mortality is not recorded after 8 days [14]. 14

Yellow phosphorus not only affects the liver, but its toxic effects are also observed in the central nervous system, including and may cause restlessness, irritability, drowsiness, lethargy, stupor, and coma owing due to liver disfunction dysfunction [15,16]; 15,16 cardiovascular toxicity with arrhythmias and hemodynamic instability [12,13], 12,13 acute tubular necrosis, and bone marrow toxicity such as

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thrombocytopenia are also observed [17]. 47 Fernandez and Canizares [18] 48 reviewed 15 cases of yellow phosphorus poisoning and found that 87% patients had some hepatic derangement after yellow phosphorus poisoning, and 27% developed fulminant hepatic failure and died. Histological analysis of the liver shows steatohepatitis and necrosis. Santos et al. [19] described three cases of white phosphorus intoxication with acute liver failure secondary to the consumption of firecrackers. In one case, liver injury improved with supportive care; in the second, the patient required liver transplantation; and the third case had a fatal outcome patient died. Similarly, Nalabothu et al. [4]4 found a 28% mortality rate in his study that was associated with the Model for End-stage Liver Disease (MELD) score, A MELD score greater than 40 was related to death, while whereas survivors presented a score lower than 12. Our patient's highest MELD score was 36. McCarron [16]<sup>16</sup> observed a mortality rate of 23-73% associated with yellow phosphorus toxicity depending on clinical manifestations; those patients with early central nervous system manifestations had a poorer prognosis. There is no antidote for phosphorus poisoning, [12,15] <sup>12,15</sup> and the only treatment is early decontamination followed by monitoring of liver function and supportive care [12,13]. 42,13 Some researchers recommend gastric lavage with 1:5000 KMnO<sub>4</sub> followed by activated charcoal and using mineral oil as a cathartic [16,17]. Fernandez et al. [18] found that NAC shows no benefits, whereas Nalabothu et al. [4]4 suggested that the early administrationuse of NAC improves outcomes for all patients with rodenticide poisonings with having liver failure, and survival rates vary with the timing of NAC administration. In their study, the survival rates was were 76%, 40%, and 23% if NAC was administered on Day 1, 2, and 3, respectively, following 40% if administered on Day 2, and 23% if administered on or after Day 3 of rodenticide ingestion [4]. However, this outcome is was confounded by an early gastric lavage in patients who were hospitalized hospitalized immediately after rodenticide ingestion. In our case, NAC was administered on Day 1 after yellow phosphorus poisoning was confirmed; however, the outcome was unfavourable. Yellow phosphorus is rapidly absorbed through the gastrointestinal mucosa, and approximately 70% accumulates in the liver within 2 to 3 hours of ingestion. It also accumulates to a lesser extent in the heart (12%), kidneys (4%), pancreas (0.4%), and brain (0.39%), and leads to damage in those organs [16]. <sup>16</sup> Histopathological changes, in our case, were mainly detected in the lungs, heart, liver, kidneys, and pancreas. No significant pathological change, other than congestion, was observed in the brain. The characteristic histopathological findings are fat infiltration, vacuolization vacuolisation, and necrosis in different organs, mainly the liver and kidney, along with focal myocarditis. Multi\_organ failure, with fulminant hepatic failure, acute tubular necrosis, and toxic myocarditis, is responsible for a fatal outcome. The toxic effect of yellow phosphorus occurs in the endoplasmic reticulum and the mitochondriamitochondrion, leading to (i) decreased synthesis of the apolipoprotein portion of very low-density lipoproteins-(VLDL), (ii) decreased production of adenosine triphosphate, and (iii) inhibition of fatty acid oxidation. This combined effect leads to fat deposition and cellular damage in different organs [20].-20

## CONCLUSION

Yellow phosphorus is a cheap and effective rodenticide. However, the number of <u>cases of</u> accidental poisonings and deaths among children and adults cannot be neglected. We agree with several other authors that the use of yellow phosphorus in rodenticides and fireworks in its currently packaged and lethal form should be banned. We have also discussed the diagnosis, and management, and <u>prognosis</u> of liver failure due to yellow phosphorus <u>poisoningand its prognosis</u>. Early and effective supportive care is <u>the key factor into</u> reducing <u>the morbidity and mortality associated with yellow phosphorus poisoning</u>.

### References

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Author Contributions: Each author is expected to have made substantial contributions to the conception or design of the work; or the acquisition, analysis, or interpretation of data; or the creation of new software used in the work; or have drafted the work or substantively revised it; AND has approved the submitted version (and version substantially edited by journal staff that involves the author's contribution to the study); AND agrees to be personally accountable for the author's own contributions and for ensuring that questions related to the accuracy or integrity of any part of the work, even ones in which the author was not personally involved, are appropriately investigated, resolved, and documented in the literature. For research articles with several authors, a short paragraph specifying their individual contributions must be provided. The following statements should be used "Conceptualization, X.X. and Y.Y.; Methodology, X.X.; Software, X.X.; Validation, X.X., Y.Y. and Z.Z.; Formal Analysis, X.X.; Investigation, X.X.; Resources, X.X.; Data Curation, X.X.; Writing - Original Draft Preparation, X.X.; Writing - Review & Editing, X.X.; Visualization, X.X.; Supervision, X.X.; Project Administration, X.X.; Funding Acquisition, Y.Y.", please turn to the CRediT taxonomy for the term explanation. For more background on CRediT, see here. "Authorship must include and be limited to those who have contributed substantially to the work. Please read th

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