Mushroom Poisoning by *Macrolepiota neomastoidea*

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There are currently over 5,000 known species of mushrooms worldwide. Only 20-25% of mushrooms have been named, and 3% of these are poisonous. More than 95% of mushroom poisoning cases occur due to difficulties associated with the identification of mushroom species. Most of the fatal mushroom poisoning cases recorded to date have been related to the *Amanita* species. Until now, a case of fatal poisoning caused by *Macrolepiota neomastoidea* (*M. neomastoidea*) has not been reported in Asia. A 57-year-old male patient was admitted to the emergency room with nausea, vomiting, diarrhea, and abdominal pain. He reported ingesting wild mushrooms with his mother and sister about 2 days ago. His mother and sister were treated with only supportive care, but he was admitted to the intensive care unit and underwent liver transplantation due to acute liver failure. We are reporting a case of fatal *M. neomastoidea* intoxication from wild mushrooms, a rare case of mushroom poisoning. (Korean J Gastroenterol 2018;71:94-97)

Key Words: Macrolepiotin; Mushrooms; Acute liver failure; Poisoning

INTRODUCTION

There are currently over 5,000 known species of mushrooms worldwide.¹ Among them, approximately 100 species are considered as toxic.² More than 95% of mushroom poisoning cases occur due to difficulties in correctly identifying the mushroom species.³ Eating wild mushrooms can cause mushroom poisoning, which can even lead to death in severe cases. We experienced a case of three family members who accidently ate *Macrolepiota neomastoidea* (*M. neomastoidea*).
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| Table 1. Laboratory Findings |
|-----------------------------|
|                           | HD 1 | HD 2 | HD 3 | HD 4 |
| AST (IU/L)                 | 2,156| 4,003| 6,273| 3,110|
| ALT (IU/L)                 | 2,155| 4,848| 7,275| 5,337|
| LDH (IU/L)                 | 2,129|     |     |     |
| PT (seconds)               | 27.1 | 65.6 | 60.7 | 91.1 |
| PT INR                     | 2.57 | 6.57 | 6.04 | 9.33 |
| Total bilirubin (mg/dL)    | 3.2  | 4.1  | 5.6  | 10.4 |
| Direct bilirubin (mg/dL)   | 2.3  | 3.0  | 3.3  | 2.3  |
| Platelet (10×3/μL)         | 252  | 231  | 56   | 22   |

HD, hospital day; AST, aspartate aminotransferase; ALT, alanine aminotransferase; LDH, lactate dehydrogenase; PT, prothrombin time; INR, international normalized ratio.

Icteric sclera were observed. His abdomen was soft with epigastric tenderness with increased bowel sounds. There were no appreciable hepatomegaly and ascites. Laboratory studies were as follows: aspartate aminotransferase, 2,156 IU/L; alanine aminotransferase, 2,555 IU/L; total bilirubin, 3.2 mg/dL; direct bilirubin, 2.3 mg/dL; blood urea nitrogen, 31 mg/dL; creatinine 1.6 mg/dL; lactate dehydrogenase, 2,129 IU/L; prothrombin time 27.1 seconds; and international normalized ratio 2.57. Hepatitis B surface antigen, anti- hepatitis C virus and immunoglobulin M hepatitis A virus were negative. Moreover, other laboratory findings, including complete blood count and alkaline phosphatase levels, were within normal range. Abdominal computed tomography revealed no abnormal findings in the liver.

He was then admitted to the general ward and rehydrated via intravenous administration with 10% dextrose fluid mixed L-aspartic acid L-ornithine 5 g and multivitamins. Complete blood count, biochemistry profile, coagulation function test, and blood gas monitoring were performed daily (Table 1). Despite the best medical supportive care, coagulation function and biochemical laboratory findings began to worsen. On hospital day 3, he started complaining of abdominal discomfort and grew lethargic. He showed worsening liver function and deep drowsy mentality with Model for End-stage Liver Disease score of 34 points. He was transferred to the intensive care unit. We decided to perform a liver transplantation. On hospital day 4, he had deceased donor liver transplantation due to acute hepatic failure. His liver weight was approximately 930 g, which is less than the weight of a normal liver. Gross finding of the liver was discoloration to reddish-green by diffuse cholestasis. The weight is approximately 930 g, which is lesser than that of normal liver.

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Fig. 3. Histopathologic findings. (A) Massive hepatocellular necrosis is present in hepatocytes. Residual viable hepatocytes show moderate macrovesicular steatosis (H&E, ×20). (B) Bile ductular proliferation (black arrows) is distinctively observed (immunohistochemistry with anti-cytokeratin18, ×20). (C, D) There is diffuse and remarkable lobular and porto-portal inflammation (H&E, ×100).

notransferase (4,298 IU/L), lactate dehydrogenase (4,882 IU/L), and prothrombin time international normalized ratio 1.34. Other laboratory findings were normal. She was fortunately improved with only medical supportive care. On hospital day 7, coagulopathy and liver function had improved and symptoms disappeared.

DISCUSSION

There are over 5,000-known species of mushrooms worldwide; however, among these, only 20-25% have been named, and only about 100 species are considered poisonous.1,2 Most patients ingesting toxic mushrooms experience no toxic effects or only experience mild-to-moderate symptoms. However, a few patients experience serious toxicity, such as hepatic failure, renal failure, and pancreatitis.4 According to the Korea Forest Service, 216 cases of mushroom intoxication and 15 deaths have been reported in Korea in the last 10 years. The major cause of death was hepatic failure from ingesting toxic mushrooms.

Such cases typically result from ingesting toxic mushrooms as a result of misidentification of the mushroom with similar morphologic features.3 In our case, the three family members mistakenly thought they were eating an edible mushroom with a similar morphology. *Macrolepiota procera* is an edible mushroom; the cap is flat, with chocolate-brown colored flakes that remain on the upper surface of the cap without any boundary between the cap and the strip (Fig. 1A). However, in *M. neomastoidea*, which is a toxic mushroom, the color changes to reddish brown when its fruit body is rubbed (Fig. 1B). This can help distinguish between the two mushrooms.

Mushroom poisoning can be divided into seven categories, depending on the type of toxin: amatoxin, gyromitrin, coprine, muscarine, ivotenic acid-muscimol, psilocybin-psilocin, and gastrointestinal irritants.5 The majority of cases of fatal toxic mushroom poisoning have been induced by amatoxin-containing mushroom, such as *Amanita, Galerina, and Lepiota* spp., and only a few cases have been caused by other toxins.6 Until now, cases of *M. neomastoidea* poisoning have rarely been reported in Asia. To the best of our knowledge, only one case was reported in Hong Kong, and it showed gastrointestinal toxicities from self-picked *M. neomastoidea* ingestion.7 However, cases of *M. neomastoidea* poisoning inducing a serious hepatic toxicity have not been reported in Asia. Therefore, our case with
M. neomastoidea-induced fatal hepatic failure requiring liver transplantation is the first in the world.

M. neomastoidea is a toxic mushroom, which is distributed throughout Korea and other East Asian countries. Mushroom poisoning by M. neomastoidea is associated with gastrointestinal symptoms, including abdominal discomfort, vomiting, and profuse diarrhea. However, whether M. neomastoidea poisoning causes liver damage and the pathogenesis of liver damage are unknown.

Macrolepiota species and Lepiota species belonging to the Agaricaceae family contain amatoxin. Amatoxin is one of the most common toxins that induce hepatic toxicity. Amatoxin, a thermoresistant toxin that enters into the systemic circulation passing through the intestinal epithelium, induces liver damage. Amatoxins interact with RNA polymerase II in the eukaryotic cells, inhibiting the process of transcription. This results in a progressive decrease in messenger RNA, causing deficient protein synthesis, hepatocyte necrosis, and liver damage. We supposed that one of the pathogenesis of hepatic injury by M. neomastoidea is amatoxin. Kim et al. reported that they isolated four chemical constituents of M. neomastoidea that were associated with cytotoxic activities against the human cells; lepiotin A, B, C, and (R)-5-hydroxypyrrolidine-2-one. Considering these findings, we assumed that M. neomastoidea can cause hepatotoxicity.

In this case, the patient's mother and sister ingested the same mushroom and were treated by only medical supportive care. However, the patient experienced acute hepatic failure and was treated by liver transplantation. The patient's mother and sister ingested less amounts of the mushroom than the patient. However, the mechanism of M. neomastoidea inducing hepatotoxicity is unknown; it can be either dose-dependent toxicity or idiosyncratic toxicity. Here, we discovered that M. neomastoidea intoxication can widely impact the liver, ranging from mildly elevated liver enzymes to acute hepatic failure requiring liver transplantation.

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