CASE REPORT

A Case of Acute Kidney Injury with Marked Hyperuricemia During Mizoribine Administration

Tomoya Nishino, Takeaki Shinzato, Yuuki Ohta, Hiroshi Yamashita, Yoko Obata, Ken Shinzato and Shigeru Kohno

Abstract

A 52-year-old woman was diagnosed with Blau syndrome and rheumatoid arthritis and was treated with prednisolone and methotrexate. Joint pain and skin ulcers were poorly controlled; therefore, mizoribine (MZ; 150 mg/day) was administered once daily from March 2011. In early July 2011, the patient was hospitalized because of acute kidney injury (AKI) and acute pancreatitis. We reasoned that AKI resulted from hyperuricemia during MZ administration because serum concentrations of uric acid (31.6 mg/dL) and MZ (trough level, 5.14 μg/mL) were markedly elevated on admission. MZ should be administered with caution because of the risk of marked hyperuricemia leading to AKI.

Key words: mizoribine, acute kidney injury, acute pancreatitis, hyperuricemia, uric acid nephropathy


Introduction

Mizoribine (MZ) is an antimetabolite, first developed in Japan. MZ is used as an immunosuppressant for collagen disease, nephrotic syndrome (NS), and organ transplantation. The immunosuppressive action of MZ results primarily from inhibition of purine metabolism, a key step in nucleic acid synthesis. Nucleic acid synthesis proceeds via 2 pathways: a de novo pathway and a salvage pathway. Lymphocytes mainly depend on the de novo pathway. MZ inhibits inosine monophosphate (IMP) dehydrogenase, which is the rate-limiting enzyme in the de novo pathway, thus inhibiting nucleic acid synthesis in lymphocytes. Other cells are less affected by MZ, because they use the salvage pathway for nucleic acid synthesis. Therefore, MZ is thought to have fewer side effects than other immunosuppressive drugs.

Hyperuricemia is a side effect of MZ treatment, and occasional cases of marked hyperuricemia leading to acute kidney injury (AKI) have been reported (2-4). Here, we report a case of hyperuricemia leading to AKI during MZ administration in a patient with Blau syndrome. Blau syndrome is juvenile sarcoidosis with an autosomal dominant pattern of inheritance and is characterized by skin rash, uveitis, and rheumatoid arthritis.

Case Report

A 52-year-old woman with Blau syndrome (juvenile sarcoidosis with an autosomal dominant pattern of inheritance and characterized by skin rash, uveitis, and rheumatoid arthritis) was treated with prednisolone (PSL) at 10 mg/day and methotrexate (MTX) at 6 mg/week from 2006 to 2011. The patient was frequently hospitalized secondary to painful, recalcitrant skin ulcers on both legs. These ulcers were likely the result of poor control of Blau syndrome. Therefore, MZ therapy was initiated in March 2011. MZ was administered at a dose of 150 mg/day once daily to control the skin ulcers and to hopefully reduce the dose of PSL. However, the patient showed poor control of the underlying Blau syndrome even after the addition of mizoribine therapy. She had arthralgia and skin ulceration with frequent remissions and exacerbations, and the inflammatory response persisted. In early July 2011, laboratory data were as follows: blood urea nitrogen (BUN), 18.6 mg/dL; creatine (Cr), 0.65 mg/dL; estimated glomerular filtration rate (eGFR), 73.9 mL/

Second Department of Internal Medicine, Nagasaki University of Medicine, Japan, Department of Nephrology, Sasebo General Hospital, Japan and Shinzato Clinic Urakami, Japan

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Correspondence to Dr. Takeaki Shinzato, takeakishinzato@yahoo.co.jp
min; and C-reactive protein (CRP), 2.4 mg/dL. Renal function was maintained, and serum concentration of MZ was 3.94 μg/mL at C2 levels. After a routine consultation in early July, she developed fever of around 38°C and painful erythema tuberculatum, which seemed to have been caused by the exacerbation of Blau syndrome. She presented with general fatigue and appetite loss, and was therefore hospitalized in mid-July 2011 for further evaluation and treatment.

The patient's height was 157 cm; weight, 61.0 kg; body temperature, 37.4°C; blood pressure, 142/90 mmHg; and pulse, 78 beats/min. Evidence of anemia or jaundice was not observed in the palpebral conjunctiva or bulbar conjunctiva, respectively. Edema was present in both legs, along with skin ulcers in several places. The laboratory data from blood and serum are given in Table 1. Laboratory evaluation on admission revealed elevated BUN (33.5 mg/dL) and Cr (4.61 mg/dL) and decreased eGFR (8.7 mL/min), indicating AKI. In addition, the uric acid (UA) level was markedly elevated (31.6 mg/dL), and the serum concentration of MZ was markedly high (5.14 μg/mL), even 24 hours after the previous oral administration (i.e., trough level). These findings suggested that MZ administration resulted in hyperuricemia and subsequent AKI. Since the patient had mild upper abdominal pain on admission, an abdominal computed tomography (CT) scan was performed to determine the cause of pain. CT findings showed no hydronephrosis or renal enlargement, but mild pancreatic enlargement was observed (Fig. 1). Laboratory data showed elevated pancreatic and hepatobiliary enzymes (amylase, 363 U/L; gamma-glutamyl transpeptidase, 222 U/L; and alkaline phosphatase, 1,127 U/L). The patient was subsequently diagnosed with comorbid acute pancreatitis.

After admission, MZ treatment was discontinued, and hemodialysis was started on day 2 in order to eliminate the UA and MZ and to treat AKI. Although oliguria was observed upon admission, urinary output began to increase on day 3. On day 6, urinary output was restored to 2,389 mL/day. Hemodialysis was discontinued after 4 sessions. After 4 weeks, renal function returned to normal.

For acute pancreatitis, the patient was treated with gabexate mesilate (600 mg/day), doripenem (0.5 g/day), and omeprazole (40 mg/day). Abdominal magnetic resonance imaging and endoscopic retrograde cholangiopancreatography were performed to investigate the cause of acute pancreatitis. No abnormalities, such as gallstones or pancreatico-biliary maljunction, were observed. The abdominal pain disappeared 2 weeks after the start of treatment. The patient recovered from AKI and acute pancreatitis, and she was discharged on day 34. The clinical course of the patient is shown in Fig. 2.

### Table 1. Laboratory Test Results on Admission

<table>
<thead>
<tr>
<th>Complete blood count</th>
<th>Blood chemistry</th>
<th>Serological test</th>
</tr>
</thead>
<tbody>
<tr>
<td>WBC 15,120 /mm³</td>
<td>TP 6.6 g/dL.</td>
<td>CRP 14.1 mg/dL.</td>
</tr>
<tr>
<td>neu 81 %</td>
<td>Alb 2.2 g/dL.</td>
<td>ANA × 40</td>
</tr>
<tr>
<td>lym 19 %</td>
<td>UA 31.6 mg/dL.</td>
<td>Anti-dsDNA Ab 6.0 U/mL</td>
</tr>
<tr>
<td>eos 0 %</td>
<td>BUN 33.5 mg/dL.</td>
<td>Anti-ssDNA Ab 23.5 U/mL</td>
</tr>
<tr>
<td>mono 0 %</td>
<td>Cr 4.61 mg/dL.</td>
<td>RF 82.7 IU/mL</td>
</tr>
<tr>
<td>baso 0 %</td>
<td>eGFR 8.7 ml/min</td>
<td>IgG 1,830 mg/dL.</td>
</tr>
<tr>
<td>RBC 321 × 10⁶ /mm³</td>
<td>Na 135 mEq/L</td>
<td>IgA 308 mg/dL.</td>
</tr>
<tr>
<td>Hb 8.5 g/dL.</td>
<td>K 4.3 mEq/L</td>
<td>IgM 237 mg/dL.</td>
</tr>
<tr>
<td>Ht 25.4 %</td>
<td>Cl 102 mEq/L</td>
<td>C3 155 mg/dL.</td>
</tr>
<tr>
<td>PLT 29.1 × 10⁴ /mm³</td>
<td>Ca 7.8 mg/dL</td>
<td>C4 41 mg/dL.</td>
</tr>
<tr>
<td>Ca 3.8 mg/dL.</td>
<td>P 3.8 mg/dL.</td>
<td>CH50 70.0 mg/dL.</td>
</tr>
<tr>
<td>T-Bil 1.0 g/dL.</td>
<td>γGTP 222 IU/L</td>
<td>MPO-ANCA &lt;10 EU</td>
</tr>
<tr>
<td>ALP 1,127 IU/L</td>
<td>γGT 187 IU/L</td>
<td>PR3-ANCA &lt;10 EU</td>
</tr>
<tr>
<td>T-cho 183 mg/dL</td>
<td>AMY 363 IU/L</td>
<td>HBs Ag &lt;2.0 S/N</td>
</tr>
<tr>
<td>TG 148 mg/dL</td>
<td>T-cho 183 mg/dL</td>
<td>HCV Ab &lt;1.00 S/C</td>
</tr>
<tr>
<td>Glu 154 mg/dL</td>
<td>Glu 154 mg/dL</td>
<td></td>
</tr>
<tr>
<td>CRP 14.1 mg/dL</td>
<td>ANA × 40</td>
<td></td>
</tr>
</tbody>
</table>

### Discussion

Here, we report a case of marked hyperuricemia leading
to AKI during administration of MZ. Hyperuricemia is a side effect of MZ and results from inhibition of IMP dehydrogenase. The incidence of hyperuricemia and AKI due to MZ has been reported to be 1.0% and 0.04%, respectively (5). MZ inhibits the production of guanosine monophosphate (GMP) from IMP and subsequently increases UA production via hypoxanthine and xanthine (6). Indeed, marked hyperuricemia has been reported in other cases of AKI after MZ administration (2-4).

UA may be involved in the development and progression of kidney diseases and may contribute to AKI. AKI is associated with increased serum UA levels as a result of both increased generation and decreased excretion of UA. In cases of MZ-induced hyperuricemia, the increased serum UA level is primarily a result of increased UA production. Increased UA levels in the blood are widely recognized to cause AKI by obstruction of renal tubules due to the formation of UA crystals within the renal tubules; this is known as acute UA nephropathy (7). Although acute UA nephropathy is typically observed as a complication of tumor lysis syndrome during antineoplastic treatment of leukemia and malignant tumors (8), it has been occasionally reported in cases of rhabdomyolysis (9) and in other conditions. Moreover, recent studies have reported that AKI due to hyperuricemia is not simply mediated by tubular cell injury but is frequently accompanied by renal vasoconstriction, microvascular injury, and a local inflammatory response (10).

MZ is excreted mainly through the kidney, and 81.1% of MZ is excreted in urine as unmetabolized drug. The excretion rate of MZ depends on renal function and is directly correlated with creatinine clearance (Ccr) (11). The half-life of MZ is 2.5 h in patients with normal renal function (Ccr >80 mL/min), 4.5 hours in patients with Ccr of 40 mL/min, 9 hours in patients with Ccr of 20 mL/min, and longer in patients with deterioration of renal function (11). Therefore, MZ dosing should be reduced in cases of decreased renal function. However, in the present case, there was no deterioration of renal function at the start of MZ administration; the patient had a Cr level of 0.55 mg/dL and an eGFR of 88.7 mL/min. Therefore, MZ was administered orally at 150 mg once daily after breakfast beginning in March 2011. Two months after beginning treatment with MZ, the patient’s serum MZ concentration was below the limit of detection; this may have been attributable to poor patient compliance.

In the present case, the MZ concentration in the blood was 3.94 μg/mL at C2 level in early July 2011. Unfortunately, serum UA levels were not measured until AKI was suspected. The patient showed poor control of the underlying Blau syndrome even after the addition of mizoribine therapy. She had arthralgia and skin ulceration with frequent remissions and exacerbations, and the inflammatory response persisted. After a routine consultation in early July 2011, she developed fever of around 38°C and painful erythema tuberculatum, which seemed to have been caused by the exacerbation of Blau syndrome. She presented with general fatigue and appetite loss, and therefore hospitalized in mid-July. Laboratory tests performed on admission revealed increased inflammatory reaction and findings that include increased white blood cell count and C-reactive protein level. Since she had already developed oliguric AKI at the time of hospitalization, we did not measure urinary uric

![Figure 2. Clinical course of the patient.](image-url)
acid levels, unfortunately. Based on the patient’s course, we presumed that the decreased renal blood flow due to inflammatory cytokine-induced increased vascular permeability, and dehydration due to pyrexia or decreased dietary intake might have led to mizoribine-induced hyperuricemia. Nevertheless, the reason for the patient not showing increased blood urea nitrogen levels was considered to be the decreased dietary intake associated with loss of appetite. Furthermore, the increased serum MZ concentration could have resulted in increased serum UA levels, rapidly causing UA nephropathy. This would have led to the deterioration of renal function. These events may have initiated a vicious cycle, thereby increasing MZ concentration in the serum. Therefore, monitoring of serum UA and MZ is necessary during MZ treatment.

In recent years, the importance of increasing the maximum drug concentration (Cmax) of MZ to adequately inhibit lymphocytes has been discussed. Although there are few reports on the effective serum peak concentrations of MZ, approximately 4-5 μg/mL of MZ is considered appropriate, as this results in a target Cmax of 3 μg/mL or higher (12, 13). Previous studies have reported that the time until maximum drug concentration (Tmax) of MZ averaged 4 hours. Therefore, it is necessary to monitor the C4 level of serum MZ. In the present case, the C2 level of serum MZ concentration had already reached 3.94 μg/mL in early July 2011, and the C4 level was estimated to be markedly higher than the target Cmax level. In fact, trough levels of MZ were very high (5.14 μg/mL) upon admission in the present case. In patients with trough levels of higher than 4 μg/mL, there is a significantly high risk of adverse effects such as thrombocytopenia, hepatic dysfunction, and mouth ulcers (14). The present patient also had comorbid acute pancreatitis of unknown origin. To date, only 2 cases of MZ-associated acute pancreatitis have been reported (15, 16). Although the precise mechanism of MZ-associated acute pancreatitis is unclear, drug-associated acute pancreatitis is thought to be due to a hypersensitivity reaction or generation of toxic metabolites. Therefore, it is important to monitor serum UA levels, renal function, pancreatic enzymes, and serum MZ concentrations (trough level and C4 level) in all patients treated with MZ.

Prevention and treatment of acute hyperuricemia and AKI are aimed at maintaining adequate hydration, reducing serum UA levels, and alkalinization of the urine, which promotes urate solubilization. Therapy for reducing UA levels includes administration of recombinant uricase, a xanthine oxidase inhibitor, and dialysis. Allopurinol is a xanthine oxidase inhibitor that blocks UA formation. Allopurinol should be used with caution because it interacts with azathioprine, resulting in bone marrow suppression. However, the use of urate-secreting drugs, such as probenecid, is contraindicated for several reasons. Urate-secreting drugs reduce the resorption of UA in the renal tubule lumen, and increase the concentration of MZ in the serum with the decline of renal function (17). Hemodialysis effectively eliminates MZ. In addition, hemodialysis can prevent the blockage of renal tubules by UA crystals. The protein-binding ratio of MZ is 1.2-5.5%, and 43% of MZ is eliminated after hemodialysis for 4 hours (18). Among 12 cases of hyperuricemia leading to AKI during MZ administration reported in Japan, 10 cases were treated with hemodialysis. In these cases, 9 of the 10 patients recovered from AKI without renal dysfunction. Similarly, in the present case, hemodialysis was performed 4 times, and there was rapid improvement in UA levels and renal function. Therefore, hemodialysis should be performed as early as possible in cases of hyperuricemia resulting in AKI during MZ administration.

In conclusion, we report a case of hyperuricemia leading to AKI during administration of MZ. During MZ administration, serum UA levels, renal function, and serum MZ concentrations (trough level and C4 level) should be closely monitored. In cases of renal dysfunction, it may be necessary to consider discontinuation of MZ and initiation of hemodialysis.

The authors state that they have no Conflict of Interest (COI).

References


