ABSTRACT

Metformin-associated lactic acidosis is a rare side effect in metformin poisoning. There is conflicting data about mortality rates changing from 3% to 83%. We aimed to discuss a case that developed lactic acidosis and acute renal failure progressing to mortality due to metformin intoxication. A 33 year-old female patient was admitted to the emergency department, with nausea and vomiting after taking 30 tablets of 1000 mg metformin. In the 2nd hour of follow-up, patient was observed to develop lactic acidosis which did not recover despite the infusion replacement treatments and hemodialysis. Following the development of respiratory failure, under mechanical ventilation, the patient developed cardiac arrest at the 48th hour of her admission. In metformin intoxications, it should be remembered that acidosis deepens very quickly and can progress with mortality despite optimal supportive therapy. More specific recommendations and further studies are required for the management and treatment of acute metformin-associated lactic acidosis.

Keywords: Metformin, lactic acidosis, intoxication, acute kidney injury

ÖZ

Metformin ile ilişkili laktik asidoz, metformin zehirlenmesinde nadir görülen bir yan etkidir. Mortalite oranı hakkında %3 ten %83’e değişen çeşitli veriler mevcuttur. Bu çalışmada metformin zehirlenmesinde başlı laktik asidoz ve akut böbrek yetmezliği gelişen ve mortalite ile sonuçlanan bir olayı tartış الأجنبية amaçladık. 33 yaşında kadın hasta, 30 adet 1000 mg metformin tablet iç-tikten sonra acil servise başvurdu ve kusma ile basvurdu. Takibinin 2. saatinde laktik asidoz geliştiği gözlemdi, infüzyon replasman tedavilerine ve hemodiyalize rağmen metabolik asidozu derinleştirdi. Solunum yetmezliği gelişen ve mekanik ventilasyon desteği verilen hastada, başvurusunun 48. saatinde kardiyak arrest gelişti. Metformin zehirlenmesinde başlı gelişen asidozun çok hızlı derinleştiği ve optimal destekleyici tedaviye rağmen mortaliteye ileriyevelileceği unutulmamalıdır. Akut metformin ilişkili laktik asidozun yönetimini ve tedavisinde daha spesifik önerileri ve daha ileri çalışmalara ihtiyaç vardır.

Anahtar kelimeler: Metformin, laktik asidoz, zehirlenme, akut böbrek hasarı
INTRODUCTION

Metformin is a reliable oral antidiabetic agent belonging to the biguanide group, which is the primary option, especially for non-insulin dependent diabetes mellitus (DM) \(^{(1)}\). Lactic acidosis is the most serious side effect of metformin use and it is the most important reason of the mortality in patients who has metformin intoxication but there are conflicting data regarding incidence of mortality. According to the data of the US poison center, 9 of 300 cases of intentional metformin overdose were mortal (3%) \(^{(2)}\). However, in one review, 5 cases of 6 deliberate metformin overdoses were mortal (83%) \(^{(3)}\).

The key management of metformin toxicity is based on ventilation and circulation support. Supportive therapy and resuscitation are essential and there is no specific antidote. Gastrointestinal decontamination should be considered in early admissions. It is important to prevent hypothermia and maintain body temperature. Bicarbonate therapy should be considered in patients with optimal ventilation and pH level below 7.0. Since sodium bicarbonate infusions alone cannot adequately correct acid-base metabolism, hemodialysis is recommended for metformin clearance and the treatment of acidosis \(^{(4)}\). Performing hemodialysis in the early period is the cornerstone of treatment to prevent mortality.

In our case report, we aimed to discuss a rare case with lactic acidosis and acute renal failure with progressing mortality despite performing early hemodialysis.

CASE

A 33-year-old female patient with known Type 2 DM, receiving metformin therapy only, admitted to the emergency department with nausea and vomiting that started 2 hours after committing suicide by taking 30 tablets of metformin 1000 mg (0.4 gr/kg total). Her general condition was good, vital parameters were as follows: blood pressure: 120/70 mmHg, heart rate: 90 per minute, respiration rate: 22 per minute. \(\text{spO}_2\): %97 and no pathological finding was found in the physical examination of the patient. Electrocardiography showed 1:1 atrioventricular nodal conduction sinus rhythm with normal QRS and QTc distances.

After the gastric lavage and activated coal (1 g/kg) therapy, hydration and symptomatic treatment were started.

No abnormality was observed in the patient’s first hemogram, liver and kidney function tests and no electrolyte imbalance was detected. Her blood gas analysis in admission was as follows: pH: 7.37, p\(\text{CO}_2\): 78.1 mmHg, HCO\(_3\): 19.6 mmol/L, lactate: 4.7 mmol/L and glucose: 186 mg/dl.

In the 2\(^{th}\) hour of her follow-up, the patient’s general condition worsened and confusion developed and her vital signs were: blood pressure: 100/67 mmHg, pulse: 100 beats/min, \(\text{spO}_2\): 99%. In the control physical examination, widespread tenderness was observed in the abdomen, and no defenses and rebounds were detected. Creatinine in control laboratory parameters was seen to increase as 2.2 mg/dl. Control arterial blood gas parameters were as follows: pH: 7.27, p\(\text{CO}_2\): 25.5 mmHg, p\(\text{O}_2\): 75 mmHg, lactate: 9 mmol/L, hCO\(_3\): 10.6 mmol/L, anion gap: 23.4, glucose: 80 mg/dL. The patient was observed to develop lactic acidosis with high anion gap due to metformin. Early hemodialysis was thought to be needed and the patient was followed up closely. Subsequent arterial blood gas parameters were as follows: pH: 6.89, p\(\text{CO}_2\): 19.5 mmHg, p\(\text{O}_2\): 75 mmHg, lactate: 18 mmol/L, hCO\(_3\): 5.6 mmol/L, anion gap: 33.4, glucose: 52 mg/dL. Treatment of hypoglycemia was started by applying intravenous 20% Dextrose 125 cc. After intravenous 100 mEq (1.5 mEq/kg) sodium bicarbonate (NaHCO\(_3\)) bolus was administered, 100 mEq NaHCO\(_3\) infusion was started. The patient, who had urinary output of 4 cc/kg/hour, was taken to hemodialysis early, as
metabolic acidosis did not improve in serial blood gas follow-ups (Figure 1) despite the infusion replacement treatments. It was observed that lactic acidosis continued to deepen in the patient who was transferred to the intensive care unit after 2 hours of hemodialysis and continuous veno-venous hemodiafiltration (CVVHF) was applied due to ongoing resistant metabolic acidosis.

Patient who developed respiratory failure in the intensive care unit and invasive mechanical ventilation treatment was initiated, cardiac arrest developed at the 48th hour of her admission. After 45 minutes of cardiopulmonary resuscitation, patient was accepted as exitus.

DISCUSSION

In our case who is a young adult patient, metformin intoxication led to acute renal failure and end up with mortality despite performing early hemodialysis.

Metformin intoxication should be suspected in a diabetic patient with high anion gap metabolic acidosis. Anamnensis and clinical findings are sufficient for diagnosis and measuring metformin level is not necessary in most cases as in our case. Although the minimum dose of metformin causing toxicity is uncertain; it has been reported that possible lethal dose concentration >50 mg/L (3). In the literature, a case has been reported in which serious side effects, especially Type B lactic acidosis, may develop after 5 gr overdose (6).

Various data are available in the literature regarding the factors predicting mortality in metformin intoxication. In a case series in which 42 patients with metformin poisoning included and mortality was seen as 48%, it was stated that the strongest predictor of mortality was liver dysfunction, especially increased prothrombin time (7). A systematic review of studies reporting acute metformin overdosing revealed that low serum pH and high serum lactate concentrations are associated with increased mortality (3). In our case, although liver dysfunction was not observed, mortality occurred with increased lactate level and decreased pH level in follow-up.

Lalau et al. (8) stated a case series that 13 cases of metformin intoxication were examined, a patient with highest lactate level as 4.2 mmol/L was treated with gastric lavage only and did not need alkalization or hemodialysis. In a case reported by Lacher et al. (9), a patient with lactate level up to 20.6 mmol/L and creatinine up to 2.4 mg/dL, was successfully treated with intravenous sodium bicarbonate replacement and hemodialysis. Although sodium bicarbonate replacement is widely used, its place in treatment is controversial.

In a literature review, it has been reported that it is vital and the most common toxicological indication for extracorporeal treatments (ECTR) in metformin poisoning (10). It has been reported that mortality rate decreases with early hemodialysis or CVVHF treatment in metformin intoxication cases resulting from suicide (11). However in another study a case treated with a short-term hemodialysis session, a rebound increase in lactate production has been reported (12). In our case, the highest lactate level was measured as 28 mmol/L; and despite sodium bicarbonate therapy, early hemodialysis and CVVHF, mortality has developed.
CONCLUSION

It should be remembered that acidosis deepens very quickly and can progress with mortality despite optimal supportive therapy. So, more specific recommendations and further studies are required for the management and treatment of acute metformin associated lactic acidosis.

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