High Anion Gap Metabolic Acidosis due to Euglycemic Diabetic Keto-acidosis Caused by Sodium-Glucose Co-transporter 2 inhibitor

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Case Report
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Case history

The case is that of 58 year-male with type 2 diabetes mellitus for 7 years, hypertension, hypercholesterolemia, who was admitted to the hospital with left lower limb cellulitis over the past 8 days. On work-up he was found to have high anion-gap metabolic acidosis (AGMA) with anion gap of 25, his lactate levels were normal (D and L-lactate). He denies overdosing with any medications and his toxicology screen for methanol, ethanol, aspirin, and ethylene glycol were negative. He has no psychiatric history of note. He denies using over the counter medications like acetaminophen. No bowel surgery could be elicited. He felt dehydrated and nauseous but otherwise fine.

His medications includes; carvedalol 25mg twice daily, hydrochlothiazide 25 mg daily, Lipitor 20 mg daily, insulin, aspirin 81 mg daily, and was started on canagliflozin 300 mg daily 4 weeks ago to control his blood sugar level and A1C.

Physical examination of the patient revealed, slightly dehydrated but well-nourished man, his vital signs; heart rate of 78 BPM and regular, BP 143/85 mmHg, temperature 98.7 F, and his oxygen saturation while breathing room air was 92%. Examination of the heart, abdomen, and chest were unremarkable. He had left lower leg cellulitis but no edema or tenderness.

His work-up including chemistry-7 which showed sodium of 142 mmol/L, potassium of 4.3 mmol/L, chloride of 102 mmol/L, bicarbonate of 13 mmol/L, BUN and creatinine of 18 mg/L and 0.78 mg/L respectively. His blood glucose level was 178 mg/L with A1C of 8.2. His serum osmolality was 312 mosm/L, and his arterial blood pH was 7.2 with a carbon dioxide in blood gas analysis (Pco2) of 32mmHg. His calculated anion gap was 25 given his normal albumin level. His investigation also showed positive ketones in the serum and urine. His urine PH was 5.5 and the urine contain >800 mg of glucose.

Case discussion

Metabolic acidosis is defined as low pH <7.3 with serum bicarbonate level of <22 mmol/L. Diabetic ketoacidosis (DKA) is one of the serious and acute complications of diabetes. It is characterized by increased total body ketone levels, metabolic acidosis, and hyperglycemia. Hyperglycemia is a key component for the diagnosis
of DKA. There is another DKA with normal blood glucose described first by Munro in 1973; where he reported 37 episodes of DKA with normal or near normal blood glucose in 17 young patients. Later on normoglycemia is defined as blood sugar <250mg/L. It is now known as Euglycemic diabetic ketoacidosis (eDKA). Burge et al have demonstrated that fasting in type 1 diabetic patients can lead to blunted hyperglycemic response which can lead to eDKA. These researchers demonstrated that dehydration can enhance the development of eDKA.

Euglycemic DKA is part of the spectrum of DKA and can be distinguished from other causes of ketoacidosis by clinical history and serum bicarbonate levels. The level of bicarbonate in starvation ketosis is usually normally more than 18 mEq/L. The absence of other causes of high anion gap metabolic acidosis like lactic acidosis, drug intoxications, and renal failure as well as the presence of ketones in the plasma and urine distinguish eDKA from other causes of AGMA. Many workers demonstrated that eDKA can occur in both type 1 and type 2 diabetes Mellitus. The most common causes that precipitated DKA are infections, inadequate insulin therapy, pancreatitis, myocardial infarction, and illicit drug use. The American Diabetic Association, defined DKA as plasma glucose of >250 mg/L, positive urinary or serum ketones, arterial PH of <7.3, serum bicarbonate <18 mEq/L, and high anion gap.

The case under discussion satisfy all the criteria of DKA except hyperglycemia. In 2013, the food and drug administration (FDA) approved many sodium-glucose co-transporter 2 inhibitor (SGLT-2 inhibitors) for management of type 2 diabetes mellitus. The use of these drugs in type-1 and 2-DM have increased the side effects and the risks of eDKA in these patients. The adverse effects of these drugs were recorded in 73 cases linked to eDKA. All patients required hospitalization and their average blood glucose level was 211mg/L. 60% of these patients have diabetes type-2. SGLT-2 inhibitors in these patients were started on average of 45 days prior to the incident. The majority of the cases had infection, dehydration, and de-escalation of insulin doses. The management of these patients were in the same line as patients with classic DKA with rehydration, insulin supplementation and treatment of the underlying precipitating cause. In some of these patients re-challenged with SGLT-2 inhibitors resulted in the development of eDKA.

Restriction of carbohydrate usage because of infection or dehydration along with de-escalation of the insulin doses because of normal blood sugars, and excretion of large amount of blood glucose in the urine as part of the effect of SGLT-2 inhibitors are suggested as the basis of the pathogenesis of eDKA. These factors increased reliance on the fat oxidation for energy with ensuing ketoacidosis.

Treatment of DKA and eDKA are the same and includes;
1. volume resuscitation, frequent assessment of cardiac function, renal function, and mental status and avoidance of fluid overload and pulmonary edema.
2. Low dose intra-venous regular insulin.
3. Correction of potassium abnormality.
4. Resolution of DKA can be assessed by the presence of 2 of the following: an anion gap <12, serum bicarbonate level >15, or a venous pH >7.3.

The important features in eDKA is the perception by the patient and the health care individual that normal blood sugar resulted in the de-escalation of the insulin dose and perpetuation of the eDKA.

Awareness that DKA can occur without high blood sugars is critical and blood gas analysis for AGMA as well as estimation of serum and urine ketones are keys for the correct diagnosis of eDKA and hence, successful management of the condition.

In conclusion: all patients with type-1 or 2 diabetes on SGLT-2 inhibitors with nausea, vomiting, shortness of breath, or malaise should have chemistry, blood gas analysis, and urine and serum ketones estimation to avert the complications of eDKA.

References


