Pituitary Apoplexy Following Severe Diabetic Ketoacidosis, With Two Uncommon Complications of Supraventricular Tachycardia and Acute Limb Ischemia in a Patient with Neglected Pituitary Adenoma and Undiagnosed Diabetes Mellitus: A Rare Clinical Association

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Abstract

Pituitary apoplexy (PA) is a clinical emergency arising from acute ischemia or hemorrhage of the pituitary gland. A small subset of pituitary adenomas present with an apoplectic crisis, with common symptoms being headache, nausea/vomiting, visual impairment, ophthalmoplegia, altered sensorium, panhypopituitarism, etc. Though diabetic ketoacidosis (DKA) is an established complication of uncontrolled diabetes mellitus, its association with PA is extremely rare. Likewise, supraventricular tachycardia (SVT) and acute limb ischemia (ALI) have rare, reported association with DKA. We present one such case of rare associations seen in our clinical practice. A 20-year-old woman was brought to our emergency room with headache, breathlessness, and altered sensorium. Clinical and biochemical evaluation revealed SVT, DKA, and right lower limb ALI. On enquiry, patient was found to be diagnosed with pituitary adenoma 2 years ago and lost to follow-up. PA was detected on neuroimaging and confirmed histopathologically. Possibility of PA presenting as DKA and its sequelae exists.

Keywords

► pituitary apoplexy
► uncontrolled diabetes
► diabetic ketoacidosis
► supraventricular tachycardia
► acute limb ischemia

Introduction

Pituitary apoplexy (PA) is a rare life-threatening condition arising from acute ischemia or hemorrhage of the pituitary gland.¹,² PA commonly occurs in the presence of pituitary adenomas, especially in nonfunctioning adenomas.¹ Incidence of PA in adenomas is underreported, as many cases may remain “clinically silent.” Common presenting
symptoms of PA include headache, nausea/vomiting, visual impairment, ophthalmoplegia, altered sensorium, panhypopituitarism, etc. Precipitating factors described for PA include hypertension, diabetes mellitus, pregnancy, radiation, etc. Relative rarity and wide spectrum of nonspecific symptoms seen in PA, makes it a diagnostic challenge.

Diabetic ketoacidosis (DKA) is a common, but serious complication of uncontrolled diabetes mellitus. Underlying pathophysiologic cause for DKA is severe insulin deficiency. Hence, DKA is more common in type 1 diabetes mellitus, either as an initial manifestation or resulting from subsequently increased insulin requirement. The spectrum of clinical presentation seen in DKA is nonspecific and vary greatly depending on the severity of the condition. Common symptoms in DKA include polyuria, polydipsia, nausea/vomiting, lethargy, breathlessness, altered sensorium, etc. Supraventricular tachycardia (SVT) and acute limb ischemia (ALI) are extremely rare complications seen in DKA. There are only a couple of published case reports on association between DKA and PA. Herein, we report the case of a young woman, known case of pituitary adenoma, presenting as DKA—complicated with SVT and ALI, secondary to PA.

**Case History**

A 20-year-old woman was rushed to the emergency room with complaints of progressively worsening headache, lethargy, breathlessness, and altered sensorium for 3 days. Due to the severity of symptoms, she was brought in a wheelchair. On arrival, she was found to be tachypneic, tachycardic (heart rate 180 beats/min), and drowsy. She had a Glasgow Coma Scale (GCS) of E3M6V4, with no gross focal deficits (quick assessment). Electrocardiogram revealed an underlying SVT as shown in [Fig. 1](#). Emergency cardiology opinion was taken, and she was immediately started on adenosine injection for cardioversion, after unsuccessful carotid massage. She also sequentially required diltiazem and amiodarone injections to get cardioverted. Meanwhile, arterial blood gas analysis revealed severe metabolic acidosis with the following readings: pH 7.083, pCO2 17.9 mm Hg, pO2 47 mm Hg, and HCO3 5.3 mmol/L. Random blood glucose was 746 mg/dL, and urine was strongly positive for ketone bodies. She was diagnosed to have DKA. Endocrinology team was consulted, and she was started on insulin infusion along with hydration. She received corrective measures for her metabolic derangements.

Within 1 hour of being in the emergency room, patient showed significant neurological deterioration. Repeat detailed neurological assessment revealed GCS of E2M5V1, with gross left-sided hemiparesis (grade 2/5) and right ophthalmoplegia. In view of her rapidly worsening neurological status and fragile cardiopulmonary condition, she was intubated—ventilated. Simultaneously, she was found to have rapidly worsening renal function leading to acute renal shutdown. After correction of intracellular compartment fluid deficit, she was started on infusion of inotropes and diuretics. Emergency computed tomography (CT) brain (plain) revealed heterogeneous, mixed density, ill-defined lesion in right suprasellar-parasellar region measuring 4.6 x 3.8 cm in the largest dimension—suggestive of pituitary adenoma with apoplexy ([Fig. 2](#)). She was then transferred from the emergency room to intensive care unit (ICU) for further management.

On detailed enquiry regarding past medical history, parents revealed that the patient had menstrual irregularities and episodes of severe headache approximately 2 years ago. She had consulted an outside physician and had undergone a magnetic resonance imaging (MRI) brain imaging as shown in [Fig. 3](#). No hormonal profiling or visual testing was done. The physician had put her on hormone stabilizing medications (no records available) for few months. This

![Fig. 1](#) A 12-lead electrocardiogram (ECG) taken at the time of arrival in the emergency room. ECG shows a heart rate of 180 beats/min, regular RR intervals, narrow QRS complexes with few p-waves buried within, and pseudo s waves (red arrow mark). These findings are highly suggestive of atrioventricular node reentrant tachycardia (AVNRT).
relieved her symptoms to a significant extent. She had no history of diabetes, though parents say she had complaints of polyuria and polydipsia in the last 2 to 3 months which was not evaluated.

Considering the emergency CT brain findings and the past medical history, patient’s hormonal profiling was done. Her cortisol (8 PM) on arrival to the hospital was 31.8 μg/dL (range 5–25 μg/dL); prolactin was 193 ng/mL (range 0–20 ng/mL); human growth hormone (GH) was 30.5 ng/mL (range 0–7 ng/mL); insulin-like growth factor 1 was 231 ng/mL (range 117–323 ng/mL); and free T3 was 4.55 pg/mL (range 1.4–4.4 pg/mL), free T4 was 2.88 ng/dL (range 0.8–2 ng/dL), and thyroid-stimulating hormone was 0.84 μIU/mL (range 0.3–5 μIU/mL). She was started on hydrocortisone injection supplementation in view of apoplectic status. Owing to the poor general condition, emergency neurosurgical intervention was put on hold.

Within 2 to 4 hours of admission to the ICU, patient’s right leg was found to be cyanosed and cold. Peripheral pulsations in the right leg were not felt from femoral level and below. Emergency Doppler study revealed thrombosis and complete obstruction of right lower limb vasculature, starting from common femoral artery and below, suggestive of ALI. Due to her then fragile status and extensive nature of the clot (no arterial flow restoration was possible as the veins were also blocked), she was not taken up for emergency vascular interventional procedure. Instead, she was started on anticoagulant therapy (low molecular weight heparin) and observed. On day 2 of admission, hemoglobin A1c done was 15.4%, suggestive of undiagnosed diabetic status.

Patient’s overall condition started improving from day 4 of admission. DKA was corrected, and acute kidney injury resolved. Her GCS improved to E4M6Vet, with spontaneous movement of right upper limb. She had residual left hemiparesis (grade ⅔) and right complete ophthalmoplegia. She got extubated on day 6 of admission. In view of the gangrenous changes that had developed in the right leg, right hip disarticulation was done on day 9 (►Fig. 4). An MRI brain (plain and contrast) was done on day 9 (►Fig. 4). An MRI brain (plain and contrast) was done at this point as patient’s kidney function normalized, and further neurosurgical plan of action was to be decided (►Fig. 5). In view of the large size and significant mass effect of the lesion, patient underwent right frontotemporal craniotomy and radical decompression of the sellar-suprasellar lesion under image guidance. She withstood the procedure well. Histopathological examination confirmed the lesion to be a nonfunctioning pituitary adenoma with apoplexy. ►Fig. 6 shows the microscopic images of the tumor. Despite the hardships of left hemiparesis and right hip disarticulation, patient is recovering well. She is in neurorehabilitation phase and on insulin therapy for newfound diabetic status. Anti-glutamic acid decarboxylase and other antibody testing are planned for defining the type of diabetes in the subsequent outpatient follow-ups.

Discussion

PA has a wide range of presentations, and hence, remains a diagnostic challenge. Common reported pathophysiological mechanisms leading to PA in adenomas are: (1) reduced blood supply due to tumor overgrowth or compression of vascular structures or other systemic factors; (2) sudden surge in blood flow due to conditions like malignant hypertension; (3) pituitary gland stimulation in pregnancy or hormonal therapy such as bromocriptine; (4) anticoagulant states; and so on.2 Though diabetes-induced
Microvasculopathy has been theoretically incriminated in the pathophysiology of PA, available clinical data does not support the same. DKA, on the other hand, has been reported to precipitate apoplectic episodes. There are only two case reports available of PA presenting as a DKA (Table 1). Jiang et al reported an apoplectic GH-secreting adenoma with DKA, while Camara-Lemarroy et al reported a nonadenomatous PA with DKA.

SVT is a broad terminology used to describe arrhythmias above or involving atrioventricular node (AVN). SVTs are a rare, reported complication of DKA. Factors precipitating SVTs in DKA could be either the severe acidosis or the accompanying electrolyte abnormalities like hypomagnesaemia or hypophosphatemia. Our case had AVN reentrant tachycardia precipitated possibly by severe acidosis, as the magnesium (2 mg/dL) and phosphorus (2.9 mg/dL) levels were normal. Vagal maneuvers and adenosine (AVN blocker) are the mainstay of treatment for SVTs in emergency conditions. Additional usage of antiarrhythmic medications is required in refractory cases. There are only three case reports available of DKA complicated by SVTs in young patients with type 1 diabetes (Table 1).

DKA has been reported to lead to a prothrombotic state, resulting rarely in devastating arterial or venous thrombotic episodes. Many postulated underlying mechanisms include dehydration and resulting hyperosmolarity/hypercoagulation state, activation of endothelium, increased platelet aggregation, impaired fibrinolysis, etc. Various reported vascular thrombotic conditions involved iliac artery, femoral artery, brachial artery, or mesenteric artery. ALI resulting from such a thrombotic episode is seldom seen (Table 1).

**Fig. 5** Contrast-enhanced magnetic resonance imaging (MRI) brain done during the present hospitalization. Lobulated dumbbell-shaped mass lesion is seen in sellar-parasellar-suprasellar region. Lesion appears mixed hypo- and hyperintense on T1-weighted images (A), heterogeneously hyperintense on T2-weighted images (B) with tiny diffuse cystic spaces at the margin of central necrotic area, and heterogeneous contrast enhancement (C, D). Size of the lesion appears significantly increased compared with previous MRI brain in Fig. 3. Findings suggestive of apoplexy in the pituitary macroadenoma.

**Fig. 6** Photomicrographs of the surgical specimen. Hematoxylin and eosin (H&E) staining showing monomorphic cell proliferation with round nuclei and chromophobe cytoplasm, indicating pituitary adenoma (red arrow), as well as intratumoral hemorrhage/necrosis (black arrow). Immunohistochemistry analysis done for various hormones show negative results, confirming nonfunctional status of the adenoma.
Our case report documents an amalgamation of the above-mentioned rare clinical associations. It highlights the worst clinical possibilities that can occur in a patient with neglected pituitary adenoma and an uncontrolled diabetes mellitus (diagnosed/undiagnosed). A watchful eye for the unfolding clinical scenario, and an appropriate early treatment can salvage even the rarest and gravest of the cases.

**Conflict of Interest**
None declared.

**References**

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**Table 1** List of available case reports on the rare clinical association between pituitary apoplexy, diabetic ketoacidosis, and acute limb ischemia

<table>
<thead>
<tr>
<th>Authors</th>
<th>Reported findings</th>
<th>Special features</th>
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<tr>
<td><strong>Pituitary apoplexy and DKA</strong></td>
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<tr>
<td>Jiang et al (2013)⁶</td>
<td>A 49-year-old man presenting with DKA was diagnosed with acromegaly and pituitary apoplexy. Due to refusal of treatment, patient had repeat episode of DKA with pituitary apoplexy after 2 months. Later died of B cell lymphoma in 3 years.</td>
<td>GH-secreting pituitary adenoma associated with DKA and apoplexy</td>
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<tr>
<td>Camara-Lemarroy et al (2016)⁴</td>
<td>A 38-year-old woman presenting with altered mental status and DKA, was found to have pituitary apoplexy. No evidence of adenoma seen.</td>
<td>DKA associated with pituitary apoplexy without underlying pituitary adenoma</td>
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<td><strong>DKA and SVT</strong></td>
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<td>Thomas et al (2007)⁸</td>
<td>Two teenage girls (13- and 14-year-old) with diagnosed type 1 diabetes mellitus status presented with DKA and SVT.</td>
<td>DKA and SVT in a diagnosed type 1 diabetes</td>
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<tr>
<td>Faruqi et al (2015)⁹</td>
<td>A 12-year-old girl with known type 1 diabetes was admitted with severe DKA and SVT.</td>
<td>DKA and SVT in a diagnosed type 1 diabetes</td>
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<tr>
<td>Finn et al (2018)¹⁰</td>
<td>An 11-year boy with undiagnosed type 1 diabetes presented with DKA and SVT.</td>
<td>DKA and SVT in an undiagnosed type 1 diabetes</td>
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<td><strong>DKA and acute limb ischemia</strong></td>
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<td>Zipser et al (2005)⁵</td>
<td>A 52-year-old man presented with acute aortoiliac and femoral artery occlusion as a complication of DKA, caused by the resulting hypercoagulable state.</td>
<td>Association between DKA and acute aortoiliac and femoral artery occlusion</td>
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<tr>
<td>Lin et al (2006)¹¹</td>
<td>A 33-year-old woman, who was a diagnosed case of diabetes, presented with DKA combined with acute brachial artery thrombosis.</td>
<td>First association reported between DKA and acute brachial artery thrombosis</td>
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Abbreviations: DKA, diabetic ketoacidosis; GH, growth hormone; SVT, supraventricular tachycardia.