Diabetic hyperosmolar nonketotic coma induced by central diabetes insipidus

Abstract

Introduction: Hyperosmolar hyperglycemic nonketotic coma (HONKC) is a rare complication of type 2 diabetes mellitus (DM). Polyuric dehydration caused by DI can lead to HONKC. One of the reasons of central diabetes insipidus (CDI) is metastatic malignancies. Most frequently breast and lung cancer and less frequently colorectal cancer can cause pituitary metastasis.

Case Presentation: A 59year old female, with a history of cervical lenf node metastasis and was being investigated for primary malignancy, admitted to emergency medicine service. She was diagnosed as HONKC with serious hypernatremia, hyperglycemia and hyperosmolarite. Although adequate hydration and normoglycemia was obtained, hypernatremia and polyuria persisted; therefore suspected diagnosis was diabetes insipidus (DI). In radiological investigation cerebral and pituitary metastatic lesions were seen, then with empirically desmopressin acetate treatment urinary output and polyuria recovered. In investigations to find primary malignancy; rectal wall was thickened and carcinoembryonic antigen levels were significantly increased. General condition was not suitable for colonoscopy and biopsy. Diagnosis was HONKC induced by DI due to pituitary metastasis of probable colorectal carcinoma.

Discussion: Polyuric dehydration caused by DI can lead to HONKC in type 2 diabetic patients. This rare togetherness should be considered in persistant hypernatremia and polyuria; although normoglycemia is provided. The most important sign of pituitary metastasis is DI. Because of this in metastatic malignencies with polyuria and polydipsia signs, pituitary metastasis should come to mind.

Keywords: diabetes mellitus, hyperosmolar hyperglycemic nonketotic coma, central diabetes insipidus, pituitary metastasis, polyuria, normoglycemia, thiazide diuretics, dehydration, enzyme inhibitor, potassium chloride, corticosteroids, hypernatremia

Case presentation

A 59year old female admitted to emergency medicine service of Canakkale State Hospital with confusion. She had comorbidities as chronic obstructive pulmonary disease (COPD) and hypertension. She had been treated with inhaler b2 agonist and angiotensine converting enzyme inhibitor therapy. One month ago she admitted to internal medicine polyclinic with polyuria and polydipsia complaints continuing for a few months. Plasma biochemistry showed: impaired fasting glucose (glucose:109mg/dl, HbA1c:6.1%), hypochromic microcytic anemia, erythrocyte sedimentation rate increased; serum electrolytes, renal and hepatic function test results were normal. Physical examination was normal except left cervical lenfadenopathy. Ultrasonography (USG) signs of lymph node was malign features; therefore excisional biopsy was performed and undifferentated carcinoma metastasis was showed (Figure 1). In investigations to find primary malignancy mammography, breast USG, abdominal USG and thorax computed tomography (CT) were normal; only in abdominal CT rectal wall was thickened.

Fever, cough, dyspnea symptoms started on the last days, oral intake was disturbed and she was taken to emergency medicine with confusion. On physical examination she was uncoincious, turgor tonus decreased, she had conjunctival paleness, no icteria, diffuse coarse rhonchus, expirium prolonged and was tachycardic. Her the calculation of body mass index (BMI) was 24kg/m² and the value was in the normal weight according to BMI classification.
Blood pressure was 100/50mmHg, heart rate was 112/min, body temperature was 38.5°C. Plasma biochemistry showed: glucose: 676 mg/dl (range: 75-100), blood urea nitrogen: 118mg/dl (range: 10-40), creatinine: 2.1mg/dl (range: 0.4-1.0), SGOT: 290IU/L (range: 10-42), SGPT: 129IU/L (range: 10-40), sodium: 172mmol/L (range: 136-144), potassium: 3.9mmol/L (range: 3.6-5.1), chloride: 130mmol/L (range: 101-111); arterial blood gas analysis: pH: 7.38 (range: 7.35-7.45), pCO2: 43.2mmHg (range: 35-45), pO2: 77mmHg (range: 70-100), HCO3: 23.4mmol/L (range: 21-26), base excess: 2.06mmol/L; urinalysis showed no keton, blood, leucocytes, protein but glucose is positive, and density: 1005g/ml; complete blood count: white blood cell: 15.8x10^3/mm^3; hemoglobin: 9.8g/dl (range: 11.7-15.5), hematocrit: 30.1% (range: 34.5-45.3), platelet: 150x10^3/mm^3 (range: 129-388), C-reactive protein: 9.8mg/dl (range: 0-0.5), erythrocyte sedimentation rate: 107mm/h (<20), Hba1c: 7% (range: 4-6). Calculated serum osmolarity was 402.5mosm/l (range: 275-295) (Table 1). The diagnosis was diabetic HONKC precipitated by pulmonary infection and the patient taken to intensive care unit.

Figure 1 solid focus of undifferentiated tumor cells inside normal lenfoid tissue and rarely seen primitive adenoid cells.

Table 1 The results of baseline laboratory tests

<table>
<thead>
<tr>
<th>Glucose</th>
<th>676mg/dl (75-100)</th>
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</thead>
<tbody>
<tr>
<td>Blood Urea Nitrogen</td>
<td>118mg/dl (10-40)</td>
</tr>
<tr>
<td>Creatinine</td>
<td>2.1mg/dl (0.4-1.0)</td>
</tr>
<tr>
<td>SGOT</td>
<td>290IU/L (10-42)</td>
</tr>
<tr>
<td>SGPT</td>
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<tr>
<td>Sodium</td>
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<td>Potassium</td>
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<tr>
<td>Chloride</td>
<td>130mmol/L (101-111)</td>
</tr>
<tr>
<td>Hba1c</td>
<td>7% (4-6)</td>
</tr>
<tr>
<td>White Blood Cell</td>
<td>15.8 x 10^3/mm^3 (4.5-11.0)</td>
</tr>
<tr>
<td>Hemoglobin</td>
<td>9.8g/dl (11.7-15.5)</td>
</tr>
<tr>
<td>Hematocrit</td>
<td>30.1% (34.5-46.3)</td>
</tr>
<tr>
<td>Platelet</td>
<td>150 x 10^3/mm^3 (129-388)</td>
</tr>
<tr>
<td>Erythrocyte sedimentation rate</td>
<td>107mm/h (&lt;20)</td>
</tr>
<tr>
<td>pH</td>
<td>7.38 (7.35-7.45)</td>
</tr>
<tr>
<td>pCO2</td>
<td>43.2mmHg (35-45)</td>
</tr>
<tr>
<td>pO2</td>
<td>77mmHg (70-100)</td>
</tr>
<tr>
<td>Base Excess</td>
<td>2.06mmol/L (±2.0)</td>
</tr>
<tr>
<td>C-reactive protein</td>
<td>9.8mg/dl (0-0.5)</td>
</tr>
<tr>
<td>Serum osmolarity</td>
<td>402.5mosm/L (275-295)</td>
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</table>

Corrected plasma sodium was 181mmol/L, and water deficit was approximately 7liters. Initially intravenous (IV) fluid replacement started with hypertonic fluids, then 0.1U/kg/h insulin infusion added and infusion rate regulated according to capillary blood glucose measurements. Potassium chloride was added to iv fluids. The patient was consulted to pulmonary disease department and antibiotic therapy started. Although blood glucose concentrations was between 100-200mg/dl with insulin infusion therapy and enough water replacement; plasma Na concentration was always above 150mmol/L and the patient remained polyuric, passing 6-8litres of dilute urine daily (urine osmolality 175-288mosm/kg). Meanwhile her confusion recovered, she was conscious, renal and hepatic function test results were normal. Primarily HONKC triggered by infection was thought in the patient.

However, suspected diagnosis was DI because of sustained high dilute urinary output although obtained normoglycemia and persistent hypernatremia resistant to fluid replacement. Sellam magnetic resonance imaging (MRI) was performed because of metastatic malignancy in history. Neurohypophysis high signal intensity loss secondary to stalk invasion in T1 weighted imaging (T1WI), stalk thickening and diffuse contrast distribution with contrasted T1WI, diffuse nodular contrasted focus due to intracerebral metastasis was showed Figure 2. General condition of the patient was not suitable for water deprivation test. Therefore, desmopressin acetate treatment was started empirically; then urinary output improved and plasma Na concentrations was between 140-145mg/dl.

Anterior pituitary hormon results was: TSH: 1.1µIU/Ml (range: 0.34-5.6), FT4: 10.5pmol/L (range: 7.9-14.4), FSH: 1.47mIU/mL (range: 7.9-14.4), LH: 0.23mIU/mL (range: 7.7-59), estradiol: 33.5pg/mL (range: 10-40), cortisole: 11µg/dL (range: 6.7-22), PRL: 41ng/dl (5-25). There was no central hypothyroidism but had secondary hypogonadism, mild hyperprolactinemia due to pituitary stalk involvement and relative hypocortisolemia. Also because of cerebral metastasis; neurosurgery department added dexamethasone for antiedema effect.

In investigations to find primary malignancy; only rectal wall thickening was shown in abdominal CT. From tumor markers carcinoembryogenic antigen levels (CEA) were >978ng/ml (range: 0-7). Colonoscopy was planned by gastroenterology department but not performed because of pulmonary dysfunction due to pneumonia and unstable general condition. Blood glucose concentrations was between 100-200mg/dl, electrolyte imbalance was recovered but unfortunately the patient died because of respiratory failure.
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Discussion

The togetherness of DI and DM in the patient has been rarely reported in Wolfram syndrome (known as DIDMOAD) as a congenital genetic disorder in pediatric age group, which has the concurrent association of CDI with type 1 DM accompanied by optic atrophy, deafness, and infantilism. But togetherness of CDI and type 2 DM is rare. Polyuric dehydration caused by DI can occur in this togetherness. In literature there are cases both caused by central and nephrogenic DI. Lithium-induced nephrogenic DI is an important side effect in the patients treated with lithium. If there are the glucose intolerance in these cases with lithium-induced nephrogenic DI, they can develop HONKC. Such cases have been described in the literature. The togetherness of CDI and type 2DM has been noted in less than 10 adult patients, some of whom had history of type 2DM before developing CDI. An exception was a patient with Klinefelter’s syndrome who had CDI for more than 5 years before presenting with hyperosmolar coma due to type 2 DM.

Furthermore, there are cases of the simultaneous development of CDI and type 2 DM reported similar to ours in the literature. Vidyarthi et al. have identified a similar case to ours in recently. They have also started desmopressin as empirically due to persistent hypernatremia and sustained polyuria in their female patient with HONKC. Then, her sodium corrected to normal with 72 hours.

Most patients who develop HONKC are elderly; but cases on the caused by DI are mostly younger as in our patient. Also frequently there is a precipitating factor. These precipitating factors are mostly infections, poor compliance with diabetic medications or drugs like corticosteroids and thiazide diuretics. In cases caused by DI; hyperosmolarity can develop without additional factors. But our patient there were many risk factors like COPD, malignancy and DI existence. Even if DI and coma was treated, immunosupression due to metastatic malignancy caused the situation to be mortal in this patient.

The initiating event in hyperosmolar hyperglycemic state is glucosuria diuresis. Glucosuria impairs the concentrating capacity of the kidney, further exacerbating water loss. The loss of more water than sodium leads to hyperosmolarity. When adequate fluid replacement and maintained normoglycemia with treatment in HONKC, renal concentrating capacity improves and thus hypernatremia and diuresis recovers. In persistent hypernatremia and sustained polyuria, additional factors should be thought. Our patient suspected diagnosis was DI because of this reason. Because of unsuitable general condition water deprivation test not performed. Recovery with empirically desmopressin treatment and sella MRI findings support the DI diagnosis.

One of the reasons of CDI is pituitary metastasis. Posterior part of the pituitary gland is the most common site of metastasis, probably due to highly rich blood supply through the hypophyseal artery. Anterior part involvement is usually seen with posterior part involvement. Most of the time this is not possible. In MRI findings high signal intensity loss of neurohypophysis and isointensity or hypointense mass is seen with T1 sequences; with T2 sequences high-intensity signal and homogeneous gadolinium involvement is seen. Our patient there was pituitary and cerebral metastasis compatible with these signs.

Breast cancer is the most common tumor metastasizing to the pituitary gland, followed by lung, prostate, renal cell and gastrointestinal cancers have also been described. In particular, CRC is a rare cause of metastasis to the pituitary gland. In two study CRC metastasis was shown to be between 2-2.4%. In this patient metastatic malignancy was the definitive diagnosis but colonoscopy and biopsy not performed because of unsuitable general condition. But rectal wall thickening in abdominal CT and high CEA levels support primary CRC. Also in lymph node biopsy there was primitive adenoid cells inside undifferentiated tumor cells. Even though CEA level is not sufficient for CRC diagnosis; it is shown to be sensitive in studies compared with healthy people.

Pituitary metastasis treatment is usually conservative and palliative. Because when pituitary metastasis is shown, the disease is usually in terminal stage. Surgical removal, radiotherapy and systemic chemotherapy can be applied in appropriate patients. In conservative treatment deficient hormonal replacement and as in our patient, steroid therapy for antiedema effect in cerebral metastasis is appropriate.

Despite all these treatments, life expectancy is to short as 6-22 months; in stalk involvement this time is 2-4 months. Our patient was died in a short time period despite conservative therapy because of additional comorbidities.

As a result in HONKC cases, seen in type 2 DM patients in younger ages without precipitating factors, additional risk factors

Figure 2 (A) In contrasted coronal and sagital T1WI stalk thickening and diffuse contrast involvement (open arrow). (B) Accompanying diffuse nodular contrasted intra cerebral metastatic focus (closed white arrow). (C) In sagital T1WI signal intensity loss secondary to neurohypophysial stalk invasion (closed white arrow).
and comorbidities should be thought. Particularly, in persistent polyuria and sustained hyponatremia DI should come to mind. Also in metastatic malignancies with polyuria and polydipsia pituitary metastasis should be suspected.

Acknowledgements

None.

Conflict of interest

Author declares that there is no conflict of interest.

References