Budd Chiari syndrome in a patient with Celiac disease: A rare entity

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ABSTRACT

Celiac disease is an immune mediated enteropathy that causes malabsorption. It is associated with a number of autoimmune diseases, however is rarely associated with Budd chiari syndrome. We present a case of a young girl who was a diagnosed case of celiac disease and had presented with abdominal distension along with pedal edema. Her initial workup was all negative while ultrasound abdomen along with CT scan abdomen had given the impression of Budd chiari syndrome. She was managed with gluten free diet, diuretics along with anticoagulants.

Key words: Celiac disease; Budd chiari syndrome; Malabsorption; Thrombolytics

INTRODUCTION

Celiac disease is one of the commonest cause of chronic malabsorption. It is an immune mediated enteropathy that is triggered by the consumption of dietary gluten in genetically prone individuals that leads to symptoms of malabsorption such as diarrhea, steatorrhea, weight loss and nutritional deficiencies.¹ The incidence of CD is considerably increased in patients having first degree family member affected with CD. The diagnosis is made when there is concordance between serological results and biopsy findings.

CD has an association with autoimmune liver diseases (primary biliary cirrhosis and autoimmune hepatitis) and an isolated hypertransaminasemia may occasional be present.² Budd- Chiari syndrome, which is characterized by the thrombosis of the hepatic veins, is a rare entity having an incidence rate of one case per 100 000 individuals.³

Celiac disease is rarely associated with Budd Chiari Syndrome and we hereby through our case report would like to highlight this association.

Case presentation

A 15 years old female girl, diagnosed case of celiac disease (based upon TTG serology and duodenal biopsy (Image 1) and compliant on gluten-free diet for the past 6 years, presented to our outpatients’ department with abdominal pain along with distension and pedal edema for past one month. Her physical examination showed pallor, mild icterus and bilateral pedal edema. Further examination
showed distended abdomen along with an enlarged and tender liver having a liver span of 16cm and a palpable spleen 4cm below the left costal margins.

Her complete blood count (CBC) revealed a hemoglobin level of 6.8g/dl, mean corpuscular volume of 63.9 fl, total leukocyte count of 9.2 x 10⁹/L, and a platelet count of 156 x10⁹/L. Peripheral blood film showed microcytic hypochromic red blood cells. Liver function tests disclosed a total bilirubin of 1.78mg/dl, alanine aminotransferase 65U/L (normal 0-37 U/L), aspartate aminotransferase 37 (0-40 U/L) and albumin 1.52g/dL. The international normalized ratio was 1.75. Viral markers were negative (HBsAg and anti HCV). Her Serum iron was low 31 µg/dL while serum Ferritin levels were within normal limits 94 µg/dL (normal 40-155 mg/dl and 11-307ug/dL). Vitamin b12 levels were normal 445 pg/mL, normal: 211–900 pg/mL).

Based upon her initial presentation, Ultrasound abdomen was done which depicted an enlarged liver having irregular margins and marked splenomegaly (16cm) along with gross ascities. Ascitic fluid analysis was done which was consistent with the findings of portal hypertension: high serum-ascites albumin gradient or gap (SAAG), low protein ascites. For further workup Serum ceruloplasmin levels were sent and were normal, while the slit-lamp examination was also negative forKayser–Fleischer ring. Autoimmune serology (anti-nuclear antibodies and anti-smooth muscle antibodies) was also negative. Thyroid profile, renal function tests were within normal limits.

Later on, a Doppler ultrasound of the liver was planned which revealed thin caliber hepatic vein, having no flow on CDI along with hypertrophy of the caudate lobe. In order to confirm our findings, we proceeded with a Contrast –enhanced computed tomography (CT) of the abdomen (Image 2), which revealed the liver to have irregular margins and appeared mottled and edematous. Caudate lobe was enlarged in size measuring 5.6x 5.2cm. Intrathepatic part of IVC was significantly thinned and there was non visualization of the hepatic veins. Multiple collaterals formation was noted at splenic hilum. A Focal filling defect of 1.1x0.9cm with surrounding calcification was noted in one of the collaterals channels, representing partial thrombus. Findings were consistent with Budd chiari syndrome.

Her Esophagastroduodenoscopy had revealed one column of grade II varices (without red sign).

Her Thrombophilia profile was unrevealing (lupus anticoagulant, protein C and protein S, antithrombotic III and factor V Leiden)

The patient was managed conservatively with anticoagulants, diuretics (spironolactone with furosemide) along with beta blockers. She was started on low salt and high protein diet along with GFD, iron and B12 supplements. On follow up after one month, patient had improvement in symptoms in terms of ascites and increase in hemoglobin.

**DISCUSSION**

This case outlines the occurrence of BCS in a diagnosed case of CD. Since she was complaint to GFD, in spite of that she developed features of BCS in the absence of any hypercoagulable state. Association of BCS with CD was first defined in 1990. Review of the literature showed that most of the cases reported were from North African Arab countries and Southern Europe. Literature review showed three cases from another region of the world (India and Argentina).

Our case is the first to be reported from Pakistan. According to the recent survey 61% of the patient of CD with BCS has no underlying thrombotic etiology as seen in our case. previously reported cases have shown that most patients have been presented in their third and fourth decade. BCS is more commonly seen in females, with some propensity to present at an advanced age than male patients. The BCS can present in acute, subacute, fulminant and chronic form in patients with CD with later being the commonest presentation similar to the patient in our case.

The most common cause of BCS is primary myeloproliferative disorders. Although nearly half of the patients have more than one causative factor accountable for the underlying hypercoagulable state. The number of theories has been suggested the association of CD
with BCS. It was proposed that the possible etiological factors are ethnic, genetic or environmental because all cases were reported from the same geographical region. Moreover, in the patients of CD associated with BCS, no specific thrombotic cause was found as in our case. It is demonstrated by several theories. Hyposplenism in CD can be responsible for the hypercoagulable state. Malabsorption of vitamin K leading to protein C, S and ant thrombin III deficiency, autoimmune vasculitis, magnesium deficiency, hyperhomocysteinemia secondary to folate deficiency or MTHFR (methylene tetrahydrofolate reductase) gene mutation are other mechanisms leading to thrombosis in BCS.

To conclude, we reported a rare association of BCS in CD from Pakistan and enlightened some factors possibly responsible for the vascular thrombosis in CD. More studies are required to know the underlying mechanism. We suggest every clinician should rule out BCS while evaluating a patient of CD of undetermined liver disease.

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