Hepatic Amyloidosis: Something That Can camouflage and Deceive our Perception!

Abstract
Amyloidosis is a multi-systemic diffusely infiltrating disease due to extracellular deposition of protein-mucopolysaccharide complexes. The type of protein deposited determines the subgroup of amyloid. Hepatic amyloidosis is a rare infiltrating disease affecting the hepatic parenchyma. A wide range of clinical presentation and atypical imaging findings delay the diagnosis of amyloidosis, while tissue biopsy demonstrating amyloid deposits is vital for a definitive diagnosis.

Keywords: B mode ultrasound, contrast CT, hepatic amyloidosis, infiltrating disease, primary secondary amyloidosis

Introduction
Amyloidosis is a multi-systemic diffusely infiltrating disease due to extracellular deposition of fibrils of protein-mucopolysaccharide complexes. Hepatic amyloidosis is a rare infiltrating disease affecting the hepatic parenchyma. A wide range of clinical presentation and atypical imaging findings delay a diagnosis of amyloidosis. Tissue biopsy demonstrating amyloid deposits is vital for a definite diagnosis.

To get a detailed insight of hepatic amyloidosis, we present a case of primary amyloidosis of liver presented to our hospital with atypical clinical and imaging findings.

Case Report
A 56-year-old male patient presented at Gastro-medicine Department with a history of abdominal distension, anorexia, weight loss, pedal edema for the past 3 months, which are progressively increasing. Had consulted an outside hospital, outside ultrasound abdomen showed hepatosplenomegaly with ascites. He came to our hospital for further evaluation and management. Blood investigations done here showed altered liver enzymes (alanine aminotransferase - 44 U/L, aspartate aminotransferase [AST] - 83 U/L, alkaline phosphatase [ALP] - 595 U/L, A/G - 2.2/3.4), creatinine of 0.8 mg/dl, prothrombin time-international normalized ratio (PT-INR) was elevated (1.86), Hb - 16.9 g/dl, platelet count - 140,000, and urine showed proteinuria (3+). Esophagastroduodenoscopy showed Grade-2 esophageal varices with gastropathy and duodenopathy. Computerized tomography (CT) abdomen with contrast [Figure 1], showed hepatomegaly with heterogeneous hepatic parenchymal enhancement with areas of hypodense areas in portal venous phase, homogenizing on delayed images (3 min); nonopacification of the right, middle, and left hepatic veins. Splenomegaly with moderate ascites. The diagnosis of Budd–Chiari syndrome was considered.

However, subsequent liver ultrasound with color Doppler [Figure 2] showed low-velocity flow in hepatic veins and portal veins with no evidence of filling defects to suggest venous thrombosis. Hepatic vein lumen was patent. Due to an elevated PT-INR (1.9) and borderline platelet count (1.4 lakhs), percutaneous liver biopsy was deferred, and transjugular liver biopsy with hepatovenous pressure gradient assessment was considered to rule out any membranous obstruction due to Budd–Chiari syndrome.

Transjugular infrahepatic inferior venacavogram and hepatic venogram showed no evidence of any filling}

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defects/stenosis in inferior vena cava/hepatic veins (right, middle, and left). Hepatovenous pressure gradient assessment showed features suggestive of portal hypertension, and subsequently, a transjugular liver biopsy was done from the right hepatic vein approach, and the sample was sent for histopathological evaluation.

Microscopy showed [Figure 3], liver tissue with distorted spatial relationship between portal tract and central vein. The parenchyma showed linear sinusoidal deposition of amorphous eosinophilic material causing atrophy of liver cell plates. The portal tracts also showed similar deposits. No significant lobular or portal inflammation. The material was congophilic on Congo red stain and showed apple green birefringence on polarizing microscopy. This congophilia persisted after pretreatment with KMnO₄. The material was weakly positive for periodic acid-Schiff. With this, a diagnosis of amyloidosis liver, possibly AL type (primary) was made.

Serum electrophoresis showed no features to suggest multiple myeloma. The patient was subsequently managed conservatively and planned to be followed up for symptomatic management.

Discussion

Amyloidosis is a multisystemic diffusely infiltrating disease due to deposition of fibrils of protein- mucopolysaccharide complexes. The type of protein deposited determines the subgroup of amyloid.

In primary amyloidosis, light chain immunoglobulin (AL) is deposited. Thus, primary amyloidosis is associated with multiple myeloma, Waldenstrom macroglobulinemia, lymphoplasmacytic lymphoma, etc. Bone marrow biopsy is required for all patients with AL amyloidosis to exclude overt multiple myeloma. In primary AL amyloidosis, the presence of light chain immunoglobulin in serum with no associated heavy chain is commoner than those associated with an IgM heavy chain which was found in around 4.4%.

In secondary amyloidosis (AA type), protein deposition happens secondary to inflammation; thus, secondary amyloidosis occurs in chronic inflammatory diseases such as rheumatoid arthritis, tuberculosis, Crohn’s disease, bronchiectasis, and chronic osteomyelitis.

Third subtype, hereditary or senile type of amyloid (ATTR) occurs due to mutant and wild type of transthyretin.

In localized or organ-limited type, amyloid material is confined to an organ, commonly the respiratory tract, skin, urinary tract, liver, etc. Localized amyloidosis is usually primary (AL type) and rarely presents as secondary (AA type). Localized amyloidosis is diagnosed primarily by excluding other systemic involvement.

In systemic type of disease or secondary amyloidosis, abnormal proteins are deposited in multiple organs, most common is the gastrointestinal tract with colon more frequently involved; other tissues such as myocardium, striated muscle, and adipose tissue, are also involved.

Amyloidosis is usually observed as a systemic form although 10%–20% is localized. Systemic amyloidosis
is progressive and fatal, death usually occurs as a result of renal or heart failure.\[^{[9]}\] In localized amyloidosis, the management is usually supportive and localized and does not require systemic therapy. Due to therapeutic and prognostic implications, distinction between the primary and secondary type of amyloidosis is clinically vital.\[^{[6]}\]

**Hepatic amyloidosis**

In hepatic amyloidosis, amyloid is deposited in the hepatic parenchyma, within the space of Disse along the sinusoids and/or along the blood vessel walls. Hepatocytes are severely compressed by the accumulation of amyloid material resulting in atrophy or near disappearance of hepatocytes. Advanced cases of amyloid infiltration, results in hepatomegaly with rubbery elastic consistency and may show “lardaceous liver” appearance on the cut surface.\[^{[9]}\]

Hepatic involvement in amyloidosis is common, although with mild clinical manifestation.\[^{[10]}\] Hepatomegaly with borderline abnormal liver function tests is more frequent findings seen with hepatic amyloidosis.\[^{[10]}\] Symptomatic presentations are due to hepatic failure, portal hypertension, and rarely due to organ rupture.\[^{[9,11,12]}\]

There has been report that 77% of hepatic amyloidosis is associated with nephrotic syndrome, congestive heart failure, peripheral neuropathy, or orthostatic hypotension.\[^{[13]}\] Other signs of hepatic amyloidosis include proteinuria (88%), elevated serum ALP (86%), abnormal serum protein electrophoresis (64%), hyposplenism on peripheral blood smear defined by the presence of Howell–Jolly bodies (62%), hepatomegaly disproportionate to abnormal liver enzymes (81%).\[^{[13]}\]

Liver function tests are not sensitive or specific, with a normal level of bilirubin and AST seen in 32% of cases.\[^{[13]}\] The median survival of patients with hepatic amyloidosis was 9 months.\[^{[14,15]}\]

**Diagnosis of hepatic amyloidosis**

Radiological findings of hepatic amyloidosis are nonspecific. Ultrasound may show heterogeneous or coarse parenchymal echotexture. Unenhanced CT shows enlarged liver with heterogeneously decreased parenchymal attenuation.\[^{[9,16]}\] Parenchymal calcifications are rarely described. On contrast enhanced CT, areas of amyloid infiltration appear as focal hypoattenuating areas, with subsequent contrast filling of these areas in delayed images. The findings are probably due to impaired blood flow as a result of amyloid infiltration of blood vessel walls and hepatocyte congestion.

Magnetic resonance imaging with oral manganese-containing contrast agent revealed several focal areas without contrast uptake by the hepatocytes and no biliary secretion in delayed images at 8 h, with no evidence of any extrahepatic biliary obstruction.\[^{[17]}\]

Definitive diagnosis requires histopathological evaluation as diagnostic imaging findings are non specific. Histologically, amyloid is recognized as a homogeneous extracellular material displaying apple green birefringence after positive Congo red staining. It shows aggregation of approximately 10 nm wide fibrils on electron microscopy, exhibits a beta-pleated sheet configuration on radiographic analysis and show resistance to proteases other than pronase.\[^{[18]}\]

**Conclusion**

- Hepatic amyloidosis, clinical presentation, and imaging manifestations are usually varied and nonspecific, which may cause a delay in diagnosis. Biopsy is always required for confirmative diagnosis.
- Cases with absent filling of hepatic veins in venous phase should always undergo a hepatic Doppler/B-mode imaging to visualize hepatic vein lumen and demonstrate venous flow. In such cases with contrast enhanced CT showing absent venous filling of hepatic veins, with hepatic Doppler/B-mode imaging showing patent hepatic vein, the possibility of underlying infiltrating pathology of hepatic parenchyma should always be considered.

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**Conflicts of interest**

There are no conflicts of interest.

**References**