DEAR EDITOR

We would like to report the case of a young lady with undiagnosed Wilson disease (WD) who presented with features of acute-on-chronic liver failure and died of its complications, which, if diagnosed in life, could have had a favorable evolution.

WD is a rare autosomal recessive disease, which is characterized by increased deposition of copper in the liver and the brain. The disease was first described in 1911 by Wilson in the monograph titled “Progressive lenticular degeneration: a familial nervous disease associated with cirrhosis of the liver.” However, it is Bramwell who has been credited to have emphasized the importance of liver pathology in the progression of WD. By the end of the 1940s, multiple authors had published the demonstrable increased deposition of copper in the liver and other organs as being associated with WD. With further understanding of the disease pathogenesis, treatment for the chelation of copper was successfully introduced with the drug penicillamine in 1956 by Walshe. This discovery of successful chelation therapy makes WD one of the most satisfying genetic diseases to be diagnosed and treated.

WD, as we understand today, is a genetic disease caused by a mutation in the gene coding for ATP7B, which is responsible for the copper-transporting ATPase. This protein is responsible for the elimination of copper into the bloodstream as well as its excretion in bile. The dysfunction of this protein, due to the mutated gene, results in an accumulation of copper within the hepatocytes. The spillage of the “free copper” results in the spread of copper into the various organs—principally the brain and cornea—resulting in oxidative damage and apoptosis of cells.

The manifestations of liver involvement have a varied spectrum depending on the stage of the disease. The earliest hepatic clinical features mirror those of non-alcoholic steatohepatitis (NASH) with histological features of micro and macrovesicular steatosis, and glycogenated nuclei. The distinction from NASH depends upon the demonstration of accumulated copper in the hepatocytes by histochemical stains. Ultrastructure shows mitochondrial changes, such as widened intercristal space, vacuolization, and increased granularity of the matrix. The intermediate stage of the disease shows histological features similar to those of autoimmune hepatitis, with the arrival of the portal and periportal inflammation composed of lymphocytes and plasma cells, which results in the destruction of the limiting plate, and parenchymal necrosis followed by bridging fibrosis. More than 50% of cases may show the presence of intra-cytoplasmic eosinophilic Mallory bodies. The cirrhotic stage may manifest as micronodular or mixed micro–macro nodular cirrhosis. The affected individual may also present with fulminant hepatic failure with histology showing features of necrosis and collapse against a background of cirrhosis.
Estimations of serum ceruloplasmin, urinary excretion of copper and liver copper content are important primary aids in the diagnosis of WD. However, the only specific diagnostic test remains the genetic analysis for the mutated ATP7B gene. However, the disease can manifest due to multiple mutations in both coding and regulatory genes in the non-coding areas. The gene spans nearly 80kb, making the tests expensive and beyond the reach of many patients.

Interpretation of simple biochemical tests have been shown to be both sensitive and fairly specific for WD. Two such indices include a ratio of alanine aminotransferase (ALT) by aspartate aminotransferase (AST), and a ratio of alkaline phosphatase (ALP) by total bilirubin (TB). An ALT/AST ratio of more than 2.2 has a sensitivity of 94% and a specificity of 86%; the ALP/TB ratio of less than 4 has a sensitivity of 94% and a specificity of 96%. These are important indices for the fact that serum ceruloplasmin—being an acute phase reactant—may be falsely high in patients with accompanying systemic inflammatory response syndrome or sepsis.

In keeping with a non-specific histomorphology of WD in the liver, the demonstration of accumulated copper remains an important tool in arriving at a diagnosis in a poor-resource setting. Orcein stain for copper-associated protein (CAP) is a useful histochemical stain. The cytoplasmic CAP on orcein stain gives a cola-colored, perinuclear, coarsely granular appearance. However, orcein positive CAP granules can be seen in chronic cholestatic disorders and are not specific for WD. The demonstration of elemental copper is possible using stains such as rubeanic acid or rhodanine. Rubeanic acid gives a greenish–black color to copper, while rhodanine stain gives a red to orange–red cytoplasmic granular appearance. However, orcein positive CAP granules can be seen in chronic cholestatic disorders and are not specific for WD. The demonstration of elemental copper is possible using stains such as rubeanic acid or rhodanine. Rubeanic acid gives a greenish–black color to copper, while rhodanine stain gives a red to orange–red cytoplasmic granular appearance. However, orcein positive CAP granules can be seen in chronic cholestatic disorders and are not specific for WD.

The hepatic parenchyma showed both micro and macrovesicular steatosis (Figure 3A). There was a marked increase in copper-associated protein as shown by the extensive cytoplasmic coarsely granular cola-colored granules on orcein stain (Figure 3C). The rhodanine stain accentuated the accumulated elemental copper within the hepatocytes in the form of red cytoplasmic granules (Figure 3D). These changes were consistent with the histopathological changes of WD.
This case is a reminder that an undiagnosed WD can present in patients, especially young females, as acute-on-chronic liver failure. In the absence of positive autoimmune and viral markers, clues should be drawn from biochemical markers, such as ALP and bilirubin in making a diagnosis. The pattern of the histopathology

Figure 1. Gross view of the liver. A – The nodular capsular surface; B – The cut surface with micro and macro nodules with intervening fibrosis.

Figure 2. Photomicrography of the liver. A – The cirrhotic pattern (H&E, 20X), which is better highlighted on Masson trichrome stain in B (20X); C – Portal tract with moderate lymphoplasmacytic cell infiltrate (H&E, 100X); D – Hepatocytes showing the presence of dense eosinophilic cytoplasmic Mallory–Denk hyaline bodies (H&E, 400X).
Liver histology and histochemistry in Wilson disease

Figure 3. Photomicrography of the liver. A – Micro and macro-vesicular steatosis (H&E, 100X); B – Intracellular and ductular cholestasis, better highlighted by emerald green staining on Fouchet stain (100X); C – Orcein stain highlighting copper-associated protein by the cytoplasmic coarsely granular cola-colored granules (100X); D – Rhodamine stain highlighting elemental copper by red cytoplasmic granules (100X).

and staining pattern of histochemical stains, such as orcein and rhodamine, demonstrated in the index case provides an overview of what to expect in a classic case of WD.

The authors retain an informed consent for the autopsy performance and the manuscript is in accordance with the Institutional Ethics committee requirements.

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