Wilson’s Disease in Bangladeshi Children: Analysis of 100 Cases

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Purpose: To evaluate clinical and laboratory profile of Wilson’s disease (WD) in children.

Methods: This cross sectional study was conducted at Bangabandhu Sheikh Mujib Medical University Hospital, Bangladesh, over a period of 3 years. One hundred consecutive children of WD between 3 to 18 years of age were evaluated.

Results: Mean age was 8.5±1.5 years. Male female ratio was 2:1. Ninety-one percent of patients were Muslim and 9% Hindu. A total of 53% cases of hepatic WD presented between 5 to 10 years of age and most of the neurologic WD manifested in 10-15 years age group. Sixty-nine children presented only with hepatic manifestations, 6 only with neurological manifestations, 14 with both hepatic and neurological manifestations, 10 children were asymptomatic and 1 patient presented with psychiatric features. WD presented as chronic liver disease (CLD) in 42%, CLD with portal hypertension in 34%, acute hepatitis in 20% and fulminant hepatic failure in 4% cases. Stigmata of CLD were found in 18% patients. Keiser-Fleischser ring was found in 76% total patients. Elevated serum transaminase was found in 85% cases, prolonged prothrombin time in 59% cases and hypoalbuminaemia in 53% cases. A total of 73% patients had low serum ceruloplasmin, basal urinary copper of >100 μg/day was found in 81% cases and urinary copper following penicillamine challenge of >1,200 μg/day was found in 92% cases.

Conclusion: Majority of studied WD children presented with hepatic manifestation of which 76% presented with CLD. Any child presented with jaundice after the age of 3 years should be investigated for WD.

Key Words: Bangladesh, Child, Wilson’s disease

INTRODUCTION

Wilson’s disease (WD) is an autosomal recessive condition affecting copper metabolism. Impaired excretion of copper through bile and decrease incorporation into ceruloplasmin causing excessive copper accumulation in different organs like liver, brain and cornea. In WD main defect remains in copper trans-
porter protein adenosine triphosphatase (ATP) 7B, a P-type ATPase resulting from gene mutation in chromosome 13. The worldwide prevalence of WD is 1 in 30,000 and the heterozygote carrier rates are about 1 in 90 persons [1]. Prevalence of WD in Bangladesh is not known.

The clinical presentation of WD varies widely. Before 10 years of age, 83% patients of WD present with hepatic manifestations and 17% with neuropsychiatric manifestations; between 10 and 18 years, 52% present with hepatic and 48% with neuropsychiatric symptoms. After the age of 18 years, 75% patients present with neuropsychiatric features and 25% patients develop liver disease [2]. The hepatic presentation also varies. Children may be asymptomatic and transaminasemia may be an incidental finding [3]. Patient may present with compensated or decompensated chronic liver disease (CLD). Sometimes splenomegaly due to portal hypertension may be the only finding [4]. Acute liver failure may be one of the presentations of WD and usually follow a bad outcome [5]. Patients provisionally diagnosed as autoimmune hepatitis and not responding to standard therapy should be investigated for WD because raised serum immunoglobulins level and autoantibodies may be found in both the conditions [6]. Coomb’s negative haemolytic anaemia is a frequent feature of WD. Hemolysis in WD may be acute or chronic, as single episode or recurrent [3]. Neuropsychiatric features usually develop in adult but may occur in children [7]. Early neurological features are behavior abnormalities, deterioration of school performance or difficulties in activities requiring hand-eye coordination [7,8]. Other neurological manifestations include deterioration of handwriting, drooling, dysarthria, dystonia, and spasticity. Seizures are infrequent in WD [9]. Common psychiatric abnormalities are depression, anxiety or even frank psychosis [10]. WD with neurological manifestation may or may not have symptomatic liver disease [11]. Occasionally patient of WD may present with renal stones, early osteoporosis, cardiomyopathy, pancreatitis, hypo parathyroidism, etc [12,13].

Prompt treatment is needed to prevent progression of neurological and hepatic damage. For this purpose, immediate diagnosis is needed. There is no single test for diagnosis of WD. Due to absence of definitive diagnostic test, identification of cases of WD is based on the combination of clinical finding, laboratory parameter and mutation analysis. Unfortunately, genetic diagnosis is not available in Bangladesh. WD disease can be diagnosed by using the scoring system developed at the 8th International Meeting on Wilson’s Disease in Leipzig, 2001 (Table 1) [14]. If proper treatment can be started before irre-

Table 1. Scoring System Developed at the 8th International Meeting on Wilson’s Disease

| Characteristic                                                                 | Score |
|--------------------------------------------------------------------------------|-------|
| KF rings (slit lamp examination)                                               |       |
| Present                                                                         | 2     |
| Absent                                                                          | 0     |
| Neuropsychiatric symptoms suggestive of WD                                     |       |
| Present                                                                         | 2     |
| Absent                                                                          | 0     |
| Coombs negative haemolytic anaemia (+high serum copper)                        |       |
| Present                                                                         | 1     |
| Absent                                                                          | 0     |
| Urinary copper (in the absence of acute hepatitis)                             |       |
| Normal                                                                          | 0     |
| 1-2×ULN                                                                         | 1     |
| >2×ULN                                                                          | 2     |
| Normal, but >5×ULN one day after challenge with 2×0.5 g D-penicillamine        | 2     |
| Liver copper quantitative                                                       |       |
| Normal                                                                          | -1    |
| Upto 5×ULN                                                                      | 1     |
| >5×ULN                                                                          | 2     |
| Rhodanine positive hepatocytes (if quantitative Cu measurement is not available)|       |
| Absent                                                                          | 0     |
| Present                                                                         | 1     |
| Serum ceruloplasmin (normal >20 mg/dL)                                         |       |
| Normal                                                                          | 0     |
| 10-20                                                                           | 1     |
| <10                                                                              | 2     |
| Mutation analysis                                                                |       |
| Disease causing mutations on both chromosomes                                  | 4     |
| Disease causing mutations on one chromosome                                    | 1     |
| No disease causing mutation detected                                            | 0     |
| Assessment of the WD-diagnosis score                                           |       |
| 4 or more: diagnosis of WD highly likely                                       |       |
| 2-3: diagnosis of WD probable, do more investigations                          |       |
| 0-1: diagnosis of WD unlikely                                                   |       |

KF: Kayser Fleischer, WD: Wilson’s disease, ULN: upper limit of normal.
versatile liver damage or neurological impairment, near normal life expectancy is possible [4].

MATERIALS AND METHODS

It was a cross sectional descriptive study. The study was conducted at the Department of Paediatric Gastroenterology and Nutrition, Bangabandhu Sheikh Mujib Medical University, Dhaka, Bangladesh, during the period of January, 2011 through December, 2013. One hundred consecutive cases of WD were studied. Patients of either sex or age range from three to eighteen years were studied. Patients presented with liver disease, seeking treatment at the Department of Paediatric Gastroenterology and Nutrition in the University Hospital, were investigated for infective, metabolic, autoimmune hepatitis and celiac disease. Detailed medical and family history was taken. Thorough physical examination was done including presence of jaundice, hepatosplenomegaly, ascites and signs of liver failure. Neurologic and psychiatric assessments were done. Slit lamp examination of eyes by an ophthalmologist was done in all cases to identify Kayser Fleischer (KF) rings and/or sunflower cataract. Complete blood count was done to see haemoglobin level and features of haemolysis. Liver function tests were done to see the extent and progression of liver disease. Serum ceruloplasmin, 24 hours urinary copper as baseline and after penicillamine challenge were done. Coomb’s test was done in all the cases of WD. All the above mentioned tests were performed using standard laboratory methods. Liver biopsy was not done because of unavailability of testing of hepatic copper in Bangladesh. Sibs of diagnosed wilsonian patients were also investigated similarly. WD was diagnosed on the basis of diagnostic scoring system proposed in Leipzig, 2001 (Table 1) [14]. Diagnosis of WD was done on the basis of presence of a score of 4 or more. Hepatic copper estimation and mutation analysis cannot be done because of unavailability of the tests in Bangladesh. CLD was diagnosed on the basis of long history of jaundice, presence of stigmata of CLD and biochemically higher aspartate aminotransferase (AST) than alanine aminotransferase (ALT) with low serum albumin. Acute hepatitis was diagnosed as acutely occurring jaundice with prodromal symptoms in previously apparent healthy individual. Acute hepatic failure was diagnosed by prolongation of prothrombin time more than 20 seconds not responded to parenteral vitamin K1, ratio of alkaline phosphatase to total serum bilirubin of < 4 and ratio of aspartate aminotransferase to ALT of > 2.2. Statistical analysis was done using SPSS Statistics ver. 18.0 (IBM Co., Armonk, NY, USA). Frequency analysis was done and results were expressed as mean, median, range and percentage.

RESULTS

During the study period 100 patients of WD were analysed. Scores of the patients of WD according to the scoring system at 8th International Meeting on WD were 4 to 5 in 64% patients, 6 to 7 in 22% and > 7 in 14% patients. Study population was 4 to 15 years with mean age 8.5 ± 1.5 years. Male female ratio was about 2:1. Ninety one percent of patients were Muslim and 9% were Hindu. Consanguinity of marriage was found in 30% of cases. Seven parents were first degree cousin. Family history of CLD was found in 15% of patients. Regarding presentation of WD 69% patients presented only with hepatic manifestation, 6% only with neurological features and 14% manifested both hepatic and neurological disease. The age of hepatic WD was between 5 to 10 years and that of neurological WD was after 10 years. Asymptomatic child was identified during family

| Age (y) | Hepatic only WD | Neurologic only WD | Hepatic with neurologic WD | Asymptomatic | Others (psychiatric) |
|---------|-----------------|--------------------|---------------------------|--------------|---------------------|
| <5      | 1               | 0                  | 0                         | 1            | 0                   |
| 5-10    | 53              | 1                  | 5                         | 9            | 0                   |
| 10-15   | 15              | 5                  | 9                         | 0            | 1                   |

Values are presented as number.

Table 2. Different type of Presentation of Wilson’s Disease (WD) in Different Age Group (n=100)
Table 3. Hepatic Presentation of Wilson’s Disease (Including Hepatic Only and Hepatic with Neurological Presentation) (n=83)

| Types of hepatic disease                  | n (%)          |
|------------------------------------------|----------------|
| Chronic liver disease                    | 35 (42.17)    |
| Chronic liver disease with portal hypertension | 28 (33.73)    |
| Acute hepatitis                          | 17 (20.48)    |
| Fulminant hepatic failure                | 3 (3.61)      |

screening (Table 2).

Hepatic presentations of WD in this study include CLD with or without portal hypertention, acute hepatitis and fulminant hepatic failure. Among the 83 hepatic cases (including hepatic only and hepatic with neurological features) CLD with or without portal hypertention is the main presentation (Table 3). Jaundice was present in 45% patients. Sixty one percent of patients had hepatomegaly and 35% had splenomegaly. Ascites was present in 38% patients. Stigmata of CLD was in only 18% patients. Common stigmata was thenar and hypothenar wasting (n=8). Spider angioma was found in two patients. Clubbing was found in only one patient. Gynaecomastia was also found in one patient. KF ring was present in 76% of the total patients. KF ring was present in 84% of the hepatic only wilsonian patients and in 90% of neurologic wilsonian patients. Asymptomatic and psychiatric patient had no KF rings. Sunflower cataract was found in 2 cases. Both patients with sunflower cataract presented with hepatic manifestation. One patient later developed pancreatitis.

Biochemical tests revealed higher serum AST level than ALT level. Prothrombin time ranges from normal to 50 seconds (Table 4).

Serum cereloplasmin was normal in 27% patient and rest of them had low serum ceruloplasmin. Baseline 24-hour urinary copper excretion was more than normal in 100% patients. Twenty-four-hour urinary copper after penicillamine challenge >200 μg/day was in 99% cases and >1,600 μg/day in 58% cases (Table 5).

In 28 cases esophageal varices was found by upper gastrointestinal tract (GIT) endoscopy. WD patient with hepatic presentations were given zinc sulphate along with penicillamine. All patients with neurological manifestation as well as asymptomatic cases were maintained on zinc.

Table 4. Liver Function Tests (n=100)

| Parameter                                      | Mean±SD       | Median (maximum-minimum) |
|-----------------------------------------------|---------------|--------------------------|
| Serum ALT (normal 15-40 U/L)                  | 108.05±102.74 | 70.0 (20-450)            |
| Serum AST (normal 15-37 U/L)                  | 124.60±72.84  | 120.0 (15-290)           |
| Serum bilirubin (normal 0.3-1.1 mg/dL)        | 3.86±3.73     | 1.90 (0.8-14)            |
| PT (control 11.8 sec)                         | 21.93±8.67    | 20.50 (13-50)            |
| Serum albumin (normal 3.5-6.5 g/dL)           | 3.09±0.88     | 3.50 (1.30-4.30)         |

SD: standard deviation, ALT: alanine aminotransferase, AST: aspartate aminotransferase, PT: prothrombin time.

Table 5. Biochemical Parameters of Copper Metabolism (n=100)

| Parameter                                      | n (%)          |
|-----------------------------------------------|----------------|
| Serum cereloplasmin (normal >20 mg/dL)        | 29 (29)        |
| <10                                           | 44 (44)        |
| 24 hour basal urinary copper excretion (normal value upto 40 μg/day) | 17 (17)        |
| 40-80                                         | 83 (83)        |
| >80                                           | 99 (99)        |
| 24 hour urinary copper after penicillamine challenge | 58 (58)    |

DISCUSSION

This study included 100 patients who were analysed clinically and biochemically. Mean age was 8.5±1.5 years. Wilson disease is rare before 5 years of age though there is report of occurance of disease at 3 years of age [15]. The youngest patient in present study was a boy of 4 years who was asymptomatic and identified during family screening. Kleine et al. [16] evaluated 28 Brazilian children, whose mean age at diagnosis was 10 years. Male are more than the female in the present study. Similar observation
was also made by some other researchers [17-20]. Merle et al. [21] found female preponderance. Some observer found no sex difference about the occurrence of WD [7]. Male preponderance may be due to the facts that parents give more importance for male child to seek medical care. Ninety-one percent of patients were Muslim and 9% Hindu. In Bangladesh, Hindus are 10% of total population. Disease frequency is almost equal in both religious groups. Consanguinity of marriage was found in 30% patients. Consanguinity of marriage is a feature of WD as because it is an autosomal recessive condition. Saito found WD to be more prevalent among the child of consanguineous parents in Japan [22].

Among the hepatic patients, 69% had only hepatic involvement and 14% had both hepatic and neurologic manifestation similar to observation of El-Karaksy et al. [17]. But a lower prevalence of 65% and 22% respectively was reported by Medici et al. [23]. High prevalence may be explained by the age of study group, where hepatic manifestations predominate in childhood. Moreover, this study was conducted at Department of Paediatric Gastroenterology and Nutrition where most of the patients come with hepatic complaints.

CLD is the main presentation of WD. Following copper deposition, hepatocytes become destroyed and result in fibrosis and cirrhosis. Portal hypertension is the result of distortion of hepatic architecture. Though WD may present with acute hepatitis, they ultimately develop chronicity [2]. WD causes hepatic failure in about 5% patients [4]. History revealed that 34% patient had gastrointestinal haemorrhage due to ruptured esophageal varices. GIT haemorrhage may also be due to coagulopathy resulted from impaired coagulation factor synthesis, qualitative or quantitative deficiency of platelets or due to peptic ulcer resulted from gastric erosion.

The prevalence of gynaecomastia in cirrhotic patients in adults has been reported in one study to be 44% [24]. Spider angioma was found in 33% cases and clubbing in 24% cases of cirrhosis [25,26]. In the present study thenar and hypothenar wasting was found in 8%, spider angioma in 2% and clubbing and gynaecomastia in 1% cases each. Spider angioma and gynaecomastia is due to impaired estradal metabolism in liver [25]. Clubbing is present in primary biliary cirrhosis and CLD with hepatopulmonary syndrome [27]. Clubbing develops due to release of platelet derived growth factor from aggregated platelets in nail bed [26]. Presence of KF rings are reliable sign in WD but it may also be found in chronic active hepatitis, primary biliary cirrhosis, chronic cholestasis and cryptogenic cirrhosis. KF rings consist of copper granules deposited in the descemet’s membrane in limbic region of cornea. Sunflower cataracts are usually associated with KF rings and they are found in anterior lens capsule with spoke radiation towards the lens periphery [28]. ALT was raised in 92% patients. Serum AST level is higher than serum ALT in this study. It is a feature of WD. Serum AST to ALT ratio >2.2 is a feature of fulminant hepatic failure due to WD [1]. ALT and AST are cytosolic enzyme and level may be normal or low in liver failure and in advanced liver disease. Ioris found normal ALT in 5.5% of patients [29]. El-Karaksy et al. [17] found median serum bilirubin of 2.9 mg/dL and median serum albumin of 2.4 gm/dL in his study on hepatic WD. In present study, median total serum bilirubin was 1.9 mg/dL, median prothrombin time 20.50 seconds and median serum albumin of 3.5 gm/dL. Liver function was not altered in a good number of patients. This may be due to the fact that these patients were picked up earlier in the disease course.

Seventy three percent patients had lower than normal serum ceruloplasmin. Ceruloplasmin is an acute phase reactant produced in liver. Low serum ceruloplasmin was found in WD, protein losing enteropathy, nephrotic syndrome, malnutrition, Menke’s disease, etc [30]. Greater than normal may be found in pregnancy, oral contraceptives usage, acute and chronic inflammation, juvenile idiopathic arthritis, Alzheimer’s disease, schizophrenia and obsessive compulsive disorder [30-32]. Dhawan et al. [33] reported normal serum ceruloplasmin in 20% patient. In our study basal 24 hour urinary copper of more than 40 μgm and post penicillamine 24 hours urinary copper more than 200 μgm (more than 5 times upper
limit normal) was found in almost all the patients. In penicillamine challenge test, 58% had 24 hours urinary copper of $>1,600 \mu g/\text{day}$. El-Karaksy et al. [17] found basal urinary copper more than $>100 \mu g/\text{day}$ in 53% patients. After penicillamine challenge it was found that 24 hours urinary copper excretion of $<1,000 \mu g/\text{day}$ in 15%, 1,000 to 1,600 $\mu g/\text{day}$ in 36.4% and $>1,600 \mu g/\text{day}$ in 48.5% patients. Limitations of this study are, hepatic copper estimation and mutation analysis could not be included in diagnostic scoring system.

WD is a treatable metabolic cause of liver disease. Majority of studied WD children presented with hepatic manifestation of which 76% presented with CLD. Any child with jaundice after 3 years of age should be evaluated for WD.

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