THERAPEUTIC RESPONSE OF WILSON’S DISEASE TO D-PENICILLAMINE IN PAEDIATRIC POPULATION: A ONE YEAR FOLLOW-UP STUDY

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Abstract:
Objectives: The objective of this study was to observe the outcome of patients treated with penicillamine.

Design: Intervention type of study

Setting: Department of Paediatrics, Dhaka Medical College Hospital

Study period: January 2007 to December 2008.

Study subjects: Sixteen diagnosed cases of Wilson’s disease as per inclusion criteria.

Intervention: D-penicillamine was started in a low dose, which was titrated gradually. The clinical and biochemical parameters were evaluated to look for the response to treatment.

Results: A total of 16 cases were included. Among them 12 were male and 4 were female. The mean (± SD) of age of the patients was 10 (± 2.34) years. Consanguinity between parents was present in 44% (n=7). The hepatic and neurological variety of WD were 56% (n=9) and 44% (n=7) respectively. The K-F ring was present in 75% (n=12/16) of WD cases. The excretion of 24 hrs urinary copper was steadily increased from discharge till second follow-up in response with increasing dose of penicillamine, thereafter the value was declining gradually till final follow-up at 1 year. Regarding outcome, 7 patients improved of which 4 were in hepatic and 3 in neurological group, 3 of hepatic WD expired and 2 developed neurological manifestations. One patients developed proteinuria while penicillamine treatment .About half of patients with WD were improved. Adequate cupriuresis occurred at three months. All the symptoms and biochemical markers WD improved gradually. No significant side effect was seen.

Key words: Wilson’s disease, penicillamine, urinary copper.

Introduction:
Wilson’s disease (WD) is a rare autosomal recessive disorder with a prevalence of 1 in 30,000 to 1 in 30,0001. It is characterized by inability of the liver to transport and store normally absorbed dietary copper, resulting in abnormal deposition of copper occurs in the liver, basal ganglia, eyes, and other tissues2.

Treatment modalities include avoidance of copper containing diet; pharmacotherapy with copper chelating agents; of these D-

Penicillamine is the oldest and most commonly used drug, which is recommended till today for the initial management of symptomatic WD3-7. Though few reports on use of zinc as treatment modality have been described but there is no universal agreement regarding its use as a single agent .Penicillamine promotes the urinary excretion of copper. The use of lower initial doses, increasing over few weeks, can increase tolerance to the drug1. Adequacy of treatment can be monitored by measuring 24-hour urinary copper excretion while on treatment1,5. Response to treatment with D-

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penicillamine has been variously described. Following treatment with penicillamine, the 24 hrs excretion of copper increases gradually at the initial phase, thereafter it begins to fall again with exhaustion of abnormally deposited copper 4,5. Though we are using penicillamine, not enough data is available in our population regarding response to penicillamine. The objective of this study was to document the outcome of WD treated with penicillamine with regard to clinical and biochemical parameters.

Materials and Methods:
This study was done in Dhaka Medical College Hospital from January 2007 to December 2008. All Paediatric patients of Wilson’s disease admitted in study place were included who fulfilled the inclusion and exclusion criteria. Diagnosis of Wilson’s disease was based on compatible clinical feature as described below plus investigation profiles. Criteria for inclusion of different types of Wilson’s disease in pediatric population (<15 yrs) were as follows 1,3,4,8,9

1) Neurological Wilson’s disease: was defined if patient had history of:
   a. Regression of acquired milestones-like progressive deterioration of school performance and of handwriting or deterioration of speech and motor function.
   b. Features of Basal ganglia lesion in the form of choreo-athetosis or dystonia with positive laboratory findings: Serum ceruloplasmin <20mg/dl, 24 hrs urinary copper (UC) >100µg/24 hrs, D-Penicillamine challenge test: 24 hrs urinary copper >1200 µg/day, presence of K-F ring on slit lamp examination.

2) Hepatic Wilson’s disease: patient with hepatitis not explainable to common causes but who fulfills compatible laboratory findings as mentioned above.

A final diagnosis was made if a patient fulfilled at least two of the following criteria 1-4 in addition to compatible clinical feature i. Presence of K-F ring on slit lamp examination
ii. Low serum ceruloplasmin level (<20mg/dl)
iii. High urinary copper excretion (>100 µg/24 hrs) iv. D-Penicillamine challenge test (UC >1200 µg/24hrs)

Exclusion criteria: Progressive neurological disease, hepatic or other clinical features compatible with WD but laboratory parameters are negative.

Study procedure:
Patients who were found eligible with regards to inclusion and exclusion criteria were enrolled after taking consent from parents. Data was collected by pre-designed structured questionnaire. After enrolment, detailed history was taken; systemic examinations and relevant laboratory investigations were done. The clinical profile and laboratory results were evaluated by a trained paediatric neurologist. Urinary copper excretion was done in all patients by “Atomic energy spectrophotometry” (normal range < 40 µg/24 hrs) from Atomic energy centre laboratory, Dhaka. Serum ceruloplasmin level was estimated by “Radial Immunodiffusion method” RN045.3 Bindarid UK, from AFIP.

Liver function tests and serologic markers for common hepatotrophic viruses like HAV, HEV, HBV, and HCV were also done to exclude other liver diseases. Routine tests like CBC, urine R/E (to detect proteinuria), serum creatinine was also done to monitor toxic effects of drugs in addition to clinically documented side effects.

Once a final diagnosis was achieved, the child was treated with D-Penicillamine (Cap Artamine 250 mg, Chandra Bhagat Pharma, Mumbai) at a dose of 5mg/kg/day then titrated with an increment of 5 mg/Kg every fortnightly to a maximum of 30mg/kg/day. In Hepatic WD increase in the dose up to 20mg/kg/day. Drugs were given before food in divided doses (after crushing granules of the capsule) along with supplementation of low copper diet and pyridoxine. Cases were then followed up for clinical and laboratory outcome by the same physician who initially assessed the cases. Neurological assessment was done for muscle power through Medical Research Council (MRC) scale(0-5,5= normal power), Grading of reflex(0-4+,4+indicates clonus) and assessment of disability by a scoring system like Barthel Index (BI index), a score of 100 BI is
continent, feeds, dresses, walks, ascends and descends himself and the severity of dystonia (Level 1-4, Level 4 = massive dystonia) was also assessed. Patients were monitored three monthly by estimating urinary copper till 1 year follow-up period to document adequate (24 hrs Urinary copper 2-5 mg/day) and maintenance cupriuresis (24 hours Urinary copper excretion 0.5-1 mg/day) along with clinical parameter. The time needed for adequate and maintenance cupriuresis was recorded. The dose and duration of drug therapy was also recorded. After completion of 1-year follow up study, patients were advised to continue the drug and to come for follow up.

Statistical analysis:
All clinical and biochemical parameters were recorded before and after the intervention. Data was edited meticulously and entered into a computer. Data was analyzed by computer-based statistical program SPSS (Statistical Package for Social Science) for Window (version 12). Results were expressed as frequency, percentage, and mean±SD. For statistical analysis paired 't' test was used for comparing means of quantitative data and Chi-square test were used for qualitative data. Differences were considered statistically significant if p<0.05.

Results:
The study was intended to observe the outcome of patients with Wilson disease following D-penicillamine treatment. The study also intended to assess the excretion of copper through the urine at different phases of treatment. A total of 16 patients were enrolled for analysis. The mean (+SD) age of the patients was 10(± 2.34) years with a range of 6-14 years. The mean (+SD) age of the hepatic and neurological WD was 7.8 yrs and 10.7 yrs with a range of 7-8 yrs and 8-12 yrs respectively. The male: female ratio was 3:1. Consanguinity between parents was present in 44% (n=7). The proportion of hepatic and neurological variety of WD was 56 % (n=9); 44% (n=7).

The K-F ring was present in 75% (n=12/16) of WD cases. It was present in all neurological WD (n=7) and 55.55% (n=5/9) of hepatic WD. The excretion of 24 hrs urinary copper was steadily increased from discharge till second follow-up in response to increasing dose of penicillamine, thereafter the value was declining gradually till final follow-up at 1 year (Table-I, II & Fig.-1).

<table>
<thead>
<tr>
<th>Disease type</th>
<th>Dose at discharge (mg/Kg)</th>
<th>Dose at Follow up (mg/Kg)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>First</td>
<td>Second</td>
</tr>
<tr>
<td>Neurological WD</td>
<td>9.29</td>
<td>30</td>
</tr>
<tr>
<td>Hepatic WD</td>
<td>5.00</td>
<td>11.88</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Disease type</th>
<th>Admission UC (µg/24 hrs)</th>
<th>Discharge UC (µg/24 hrs)</th>
<th>Follow up UC (µg / 24hrs)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1st</td>
<td>2nd</td>
<td>3rd</td>
</tr>
<tr>
<td>Neurological WD</td>
<td>287.14</td>
<td>1724.49</td>
<td>3391.0</td>
</tr>
<tr>
<td>Drug dose (mg/kg)</td>
<td>nil</td>
<td>9.29</td>
<td>30</td>
</tr>
<tr>
<td>Hepatic WD</td>
<td>360</td>
<td>1552.89</td>
<td>2263.0</td>
</tr>
<tr>
<td>Drug dose (mg/kg)</td>
<td>nil</td>
<td>5</td>
<td>11.8</td>
</tr>
</tbody>
</table>
Compared to admission, the 24 hrs urinary copper excretions were increased progressively in subsequent follow-up visits in response to penicillamine therapy \((p < 0.05)\) - Table-III. At first follow up, all of neurological Wilson disease and 5 (62.5%) of hepatic Wilson disease patients had adequate cupriuresis (urinary copper excretion 2-5 mg/day), none of the patient achieved a state of maintenance cupriuresis (urinary copper excretion 0.5 – 1 mg/day) at 1yr final follow-up. The concentration of urinary copper excretion increased initially following chelation with D-Penicillamine, thereafter attained a steady state before being declined gradually in both the groups on subsequent follow up (Fig-1).

The mean value at admission of both serum bilirubin (10.14 mg/dl) and SGPT (80.56 IU/L) was decreased gradually to final value of 3.2 mg/dl and 39.60 IU/L respectively in hepatic Wilson’s disease. The INR value was also decreased significantly in subsequent follow-up compared to admission \((p < 0.05)\). For neurological WD there was no change in the grading of muscle power but slight improvement \((p=0.05)\) in the grading of reflexes was observed, severity of dystonia and disability score (BI) showed a trend of improvement though statistically not significant till final follow-up.

#### Table-III

<table>
<thead>
<tr>
<th>24 hrs UC</th>
<th>Neurological Wilson’s</th>
<th>Hepatic Wilson’s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean drug dose mg/kg</td>
<td>24 hrs UC</td>
<td>Mean drug dose mg/kg</td>
</tr>
<tr>
<td>t</td>
<td>df</td>
<td>p</td>
</tr>
<tr>
<td>---</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>A</td>
<td>9.29</td>
<td>- 5.766</td>
</tr>
<tr>
<td>B</td>
<td>30</td>
<td>- 5.451</td>
</tr>
<tr>
<td>C</td>
<td>30</td>
<td>- 8.289</td>
</tr>
<tr>
<td>D</td>
<td>30</td>
<td>- 6.894</td>
</tr>
<tr>
<td>E</td>
<td>30</td>
<td>- 5.632</td>
</tr>
</tbody>
</table>

**Fig.-1:** Line chart showing 24-hour urinary copper excretion at admission, discharge and subsequent follow up till 1 yr.

**Fig.-2:** Line chart showing disability score of neurological WD at admission, discharge and different follow-up.
Table 4 shows that among the hepatic WD, 4 patients improved, 3 expired and 2 developed neurological manifestations. In neurological WD, 3 patients improved and 4 cases remained unchanged.

**Discussion:**
The study included the patients with Wilson’s disease and intended to observe the outcome of patients treated with D-Penicillamine. The mean (±SD) age of the patients was 10(± 2.34) years with a range of 6-14 years. This finding is similar to the study conducted by Yuce et al (2003) who found the age range from 6-15 years. The mean (±SD) age of the hepatic and neurological WD was 7.8 yrs and 10.7 yrs with a range of 7-8 yrs and 8-12 yrs respectively. The mean age of presentation of hepatic WD at 6.8 yrs and neurological WD at about 8 yrs has also been reported by Kalra et al (2000) from India.

The male female ratio of the patients was 3:1. The sex distribution is quite similar to the findings of Shakya and Agrawal (2004) who got 73.7% patients being male. 43.75% patients have their parents with consanguineous marriage. Parental consanguinity was noted in 12.5%, among hepatic WD in another study from Bangladesh and also from Turkey where consanguinity was as high as 70 % due to intra-family marriage which also seems to be likely in the present series.

There is a great deal of variability in the clinical presentation, typically a hepatic WD presents in the first decade of life, neurological Wilson’s generally due to extrapyramidal involvement and psychiatric symptoms usually presents later. In broad terms, patients can present acutely with liver failure, haemolysis or both or more chronically with liver disease, neurological disease or both. In the present study hepatic WD was more common (56%, n=9) than neurological WD (44%, n=7) which is similar to a study from India where out of 124 paediatric patients, hepatic WD was observed in 54%(n=67) cases.

K-F rings are usually present in 95% of children with neurological symptoms, 50-60% of children without neurological symptoms and in 10% of asymptomatic siblings with WD. K-F ring was present in all patients of neurological WD disease and in 55.56% patients of hepatic WD disease which corroborates with above report. Yuce et al (2003) also demonstrated the presence of K-F ring in all patients with neurological manifestation and in 58% patients with only hepatic presentation. Shakya and Agrawal’s (2004) observation also supports the present report. They reported K-F ring in 89.5% of patients.

The mean (± SD) of 24 hours urinary copper at admission was 328.13 (± 169.69) µg. This value is supported by Roberts and Schilsky (2003). They stated that excretion of urinary copper greater that 100 µg / 24 hours is a diagnostic marker of Wilson’s disease. The 24 hours urinary copper excretion with a range of 112-2800 µg/d was also observed by Yuce et al. After giving D-Penicillamine the mean (±SD) urinary excretion of Cu increased to 1613.94 (±282.93) µg at discharge. It was then progressively increased to a value of 2759-2790 µg/24 hrs at 2nd and 3rd follow-up, to achieve a steady state in between 3-6 months after starting penicillamine. The 24 hours excretion thereafter declined gradually to a value of 1501.50 (±758.06) µg at 1 yr -final follow-up. This observation supports the view that following treatment with penicillamine, the 24 hrs excretion of copper increases gradually at the initial phase, thereafter it reaches a steady state until it begins to fall again with

<table>
<thead>
<tr>
<th>Type of disease</th>
<th>Improved</th>
<th>Expired</th>
<th>Unchanged</th>
<th>Developed new problem</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hepatic WD</td>
<td>4</td>
<td>3</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Neurological WD</td>
<td>3</td>
<td>0</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td><strong>7</strong></td>
<td><strong>3</strong></td>
<td><strong>4</strong></td>
<td><strong>2</strong></td>
</tr>
</tbody>
</table>
exhaustion of abnormally deposited copper. The majority of the patient achieved adequate cupriuresis at 3 months following penicillamine that was maintained till 6 months, thereafter started declining. None of the patient of the present study achieved maintenance cupriuresis at 1 year follow-up, documenting that more time is required to achieve that state. An improvement in the clinical features following 2-3 months of treatment, continuing over a period of 1-2 years along with an increase in 24 hours UC has also been reported in the literature.

A marked increase in the urinary copper excretion after one week of penicillamine treatment was also described by Karim et al. This finding is also supported by Bertrand et al. They stated that following D-Penicillamine therapy urinary copper excretion initially increased and thereafter gradually decreased to approximately 50% of initial values.

The D-Penicillamine therapy in hepatic WD of the present series caused gradual decrease in serum bilirubin from initial value of 10.1 (±9.64) mg/dl to a value of 3.2 (±3.36) mg/dl at final follow up. Similarly SGPT decreased from admission value of 80.56 (±73.86) IU/L to final follow up value of 39.60 (±39.66) IU/L and significant reduction in INR from admission value of 1.64 (±0.6) sec to a final value of 1.2 (±0.45) sec. These findings are supported by Roberts and Schilsky (2003), Durand et al (2001), and Ala et al (2007). Roberts and Schilsky (2003) described that improvement in synthetic function of liver and clinical sign like Jaundice occurs during the first 2 to 6 months of treatment, but further recovery can occur during the first year of treatment. Durand et al (2001) experienced that early administration of D-penicillamine in patients with severe hepatic insufficiency was associated with survival without liver transplantation. Ala et al (2007) stated that high bilirubin, prolonged prothrombin time usually improved with D-penicillamine treatment in patients of WD with severe liver disease.

In neurological WD clinical improvement was not so evident in early stage but definite improvement was noticed in biochemical parameters as reflected by progressive increase in excretion of urinary copper, increasing trend of improvement in disability score and other clinical parameters on follow up.

Following treatment with penicillamine, though the disability score increased gradually but it was not statistically significant. Similarly dystonia and muscle power improved but did not reach to the significant level. The non significant neurological improvement was probably related to the irreversible changes in basal ganglia as part of pathogenesis. These findings were supported by Kalra et al (2000) and Ala et al (2007). In a study with 25 Indian children of WD treated with D-penicillamine, zinc, pyridoxine and low copper diet, Kalra et al found improvement in majority, residual dysarthria in seven, prolonged persistence of K-F ring in 15, and complications like renal tubular acidosis with osteopenia in one. Ala et al (2007) described that patients with neurological WD showed gradual clinical and cerebral MRI improvement with penicillamine treatment.

One of our hepatic WD patients also had joint pain, difficulty in walking. X-ray of wrist and ankle joints showed widening of epiphysis of both wrist and ankle joints. After treatment with dicalcitol, calcium and penicillamine, joints became normal in final x-ray follow-up. This observation was supported by Roberts and Schilsky (2003). They reported that patients with WD may present with important extra-hepatic manifestation like premature osteoporosis and arthritis.

Side effects:
One of our patients developed proteinuria as a side effect of penicillamine in which it was replaced by zinc. Another two hepatic WD developed neurological symptoms like dystonia, dysarthria while on treatment. All these recognized side effects are supported by Roberts & Schilsky. Roberts and Schilsky stated that penicillamine is associated with numerous side effects. Early sensitivity reaction includes fever, cutaneous eruption, neutropenia, thrombocytopenia and proteinuria and late
reaction include nephrotoxicity heralded by proteinuria or the appearance of other cellular elements in the urine for which discontinuation of penicillamine should be immediate.

Final outcome:
Out of 16 cases, 7 improved of which 4 in hepatic and 3 in neurological group, 3 hepatic WD expired and 2 developed neurological manifestations. In neurological WD 4 cases remained unchanged till study period though 24 hours UC excretion was significant in all cases. The improvement in majority of the cases treated with D-penicillamine had also been reported by Kalra et al, Shakya et al and Durand et al 2,14,17.

Conclusion:
About half of patients with WD treated by D-penicillamine were improved. Adequate cupriuresis occurred at three months; none achieved maintenance cupriuresis till one year after administration of the drug. All the symptoms and biochemical markers of WD improved gradually. No significant side effect was observed in the majority.

References: