The Clinical Utility of a Low Serum Ceruloplasmin Measurement in the Diagnosis of Wilson Disease

Abstract:

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The first step in screening for potential Wilson disease is serum ceruloplasmin testing, whereby a level of less than 0.2g/L is suggestive of the disease. We aimed to determine what proportion of an Irish population had a low ceruloplasmin level, whether low measurements were appropriately followed-up and what were the clinical outcomes. We conducted a retrospective review of all serum ceruloplasmin measurements between August 2003 and October 2009 in a large tertiary referral centre in Southern Ireland. Clinical data, serum ceruloplasmin, liver function tests, urinary copper and liver biopsy reports were all recorded where available. 1573 patients had a serum ceruloplasmin measurement during the 7-year study period. 98 patients (6.1%) had a ceruloplasmin level <0.2g/L and of these only 3 patients had Wilson disease. There was only 1 new diagnosis. Only 27 patients (28.1%) had some form of confirmatory testing performed. In our centre's experience, the positive predictive value of a significantly low ceruloplasmin level is 11.1% (95% CI 2.91-30.3%). In practice a low serum ceruloplasmin measurement is often not followed by appropriate confirmatory testing. Measuring serum ceruloplasmin as a singular diagnostic test for Wilson disease or as part of the battery of unselected liver screening tests is inappropriate and low-yield.

Introduction

Wilson disease (WD) is a rare autosomal recessive defect in hepatocellular copper transport found in 3 out of 100,000 people. It can lead to chronic copper deposition in the liver, brain and other tissues resulting in hepatotoxicity and neuropsychiatric sequelae. The American Association for the Study of Liver Disease (AASLD) recommends screening for WD in any individual aged between 3 and 55 years with liver abnormalities of uncertain cause, especially those with co-morbid unexplained neurological disorders. The first step in screening for potential WD is serum ceruloplasmin measurement, as approximately 95 to 99 percent of patients with WD have low serum ceruloplasmin levels. However, low ceruloplasmin is not specific for WD; it can result from malabsorption, other liver diseases, protein-losing enteropathies, acquired copper deficiency, and hereditary aceruloplasminemia. Ceruloplasmin is also an acute phase reactant and may be elevated in inflammatory states including WD patients with active hepatitis. Hyper-estrogenic states including pregnancy or use of the oral contraceptive pill can also increase ceruloplasmin levels as ceruloplasmin mRNA has an estrogen responsive upstream region for its transcription. Ceruloplasmin levels of less than 0.2g/L have been shown in one study to have a sensitivity of over 98%, specificity of over 55% and positive predictive value of over 48% for the diagnosis of WD on genotype-verified patients.

However, other research suggests the positive predictive value of ceruloplasmin <0.2g/L when used alone in patients with liver dysfunction may be as low as 5.9%. For this reason, the guidelines recommend a number of confirmatory tests including a slit-lamp examination for detection of Kayser-Fleischer (KF) rings and 24-hour urinary copper estimation. Additional investigations may be required for those with indeterminate results including a liver biopsy to determine the hepatic copper concentration or molecular testing for ATP7B mutations. The European Association for the Study of the Liver (EASL) clinical practice guidelines also recognizes the limitations of serum ceruloplasmin measurement as a single test and recommends use of a combination of tests that reflect disturbed copper metabolism with a diagnostic scoring system based on their results. Our study assessed the incidence of low ceruloplasmin levels in the population and the clinical indications for testing. We determined whether further diagnostic testing was performed and what was the clinical significance of these results.

Methods

We conducted a retrospective review of all ceruloplasmin measurements at the Cork University Hospital clinical laboratory that serves a large primary care network, specialty clinics, and an 800-bed tertiary neurological and hepatological referral centre between August 2003 and October 2009. The time period was chosen to include the maximum data available. The study was approved by the hospital ethics committee. Clinical data, serum ceruloplasmin, liver function tests, urinary copper and liver biopsy reports were all recorded where available. Nephelometry was the technique used by the referral laboratory for measurement of serum ceruloplasmin during the study period. We treated ceruloplasmin values less than 0.2g/L as potentially positive for WD in accordance with published guidelines. A diagnosis of WD was determined on the basis of the results of confirmatory testing (urine and hepatic copper, ophthalmologic examinations) and medical record documentation. Genetic testing results were recorded where available. WD heterozygotes were defined as those with only one disease-causing mutation in the absence of other supportive features. Data was analysed using SPSS version 21.
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Results

Demographics of the tested population

Our laboratory in Cork city, Ireland serves as a supra-regional centre for a total population of 1.1 million people. There were 1573 patients in the laboratory database who had a serum ceruloplasmin measurement during the 7-year study period. Table 1 outlines the demographic details and referral source by clinical speciality. A total of 476 serum ceruloplasmin levels were ordered in patients aged less than 3 years or greater than 55 years, a group outside AASLD recommendations. Please see Figure 1 for the range of ceruloplasmin levels in the entire population. The median ceruloplasmin level for all patients was 0.28 g/L (5th centile = 0.18, 95th centile = 0.5 g/L) with a male to female ratio of 72:28% and 64.2% of the population were aged less than 20 years. The reference range in the median for males and females at 0.27/Lg and 0.3/Lg respectively (p<0.05). There was a weak positive correlation between age and ceruloplasmin level with lower ages associated with lower ceruloplasmin level, r = 0.124, n = 1573, p < 0.005.

Positive ceruloplasmin results

96 patients (male:female = 72:28%) of the entire cohort (6.1%) had a ceruloplasmin level <0.2g/L (see Figure 1). The most common indications for testing were the presence of liver function test abnormalities (61.5%), a movement disorder (18.8%) and psychiatric disturbances (7.3%). 46.6% of the group were subsequently given a definitive liver disease diagnosis, the most common of which was alcoholic liver disease (14.6%) followed by non-alcoholic fatty liver disease (9.5%). Less than 5% (3.1%) had WD. Two of these patients had been previously diagnosed and were re-tested for disease monitoring. One patient was newly diagnosed with WD during the study period. His elevated 24-hour urinary copper excretion, Kayser-Fleisher rings along with low serum ceruloplasmin level confirmed the diagnosis. All 3 WD patients had serum ceruloplasmin levels <0.10 g/L. In addition, 1 individual presenting with tremor was verified to have WD carrier with a single ATP7B mutation only. His genetic analysis was not consistent with a compound heterozygote and he never developed any further clinical symptoms.

Confirmatory testing for Wilson disease

After a positive/suggestive ceruloplasmin result, 27/96 patients (28.1%) had some form of confirmatory testing for WD performed. 14 patients (14.5%) were examined ophthalmologically for Kayser-Fleisher rings. 10 patients (10.4%) had a liver biopsy for hepatic copper level quantification. 23 patients (24%) had urinary copper quantification. The mean follow-up period from the time of initial testing was 51.4 months (SD 30.3 months). Non-neurology/gastroenterology subspecialties were significantly less likely to do follow-up investigations (p<0.01). Only 4 of the 27 patients (14.8%) who had further testing were under the care of the non-neurology/gastroenterology subspecialties.

Ceruloplasmin as a screening diagnostic test

In our centre experience, the positive predictive value (PPV) of a ceruloplasmin level <0.2g/L is 11.1% (95% CI 2.91-30.3%). The false positive rate is 88.9% (95% CI 69.7-97.1%). At a hypothetical lower ceruloplasmin cut-off of 0.14 g/L, we observed a PPV of 22.2% (95% CI 12.6-32.8%). This results in a large increase in sensitivity (all 3 WD cases had a serum ceruloplasmin <0.05g/L). There was a statistically significant relationship between gender, age, and likelihood of a positive ceruloplasmin test result. The mean age of those with a positive level was 36.5 years compared to 45.8 years in those without (p=0.03). 10% of the tested male population had a low level versus 5.5% of the female population (p=0.01). There was a three-fold increase in frequency of ceruloplasmin testing per year over the 7 years studied comparing the tests done in 2003-04 (109) to 2008-2009 (362).

Discussion

Measurement of ceruloplasmin is often prompted in the evaluation and management of the patient with liver enzyme elevation observed on routine screening patterns, we found poor adherence to the AASLD guidelines. A significant proportion of the tested population was outside of the age-range recommended by the AASLD. It is also clear from our data that a large number of patients with low ceruloplasmin levels in 5023 patients, the number needed to test for late-onset Wilson disease (>40 years of age) to result in 1 new diagnosis was 2847. In many cases, the liver abnormalities were not of uncertain aetiology and more common diseases were not yet ruled out. Several of those with low ceruloplasmin levels had chronic hepatitis or alcohol-induced liver disease conditions associated with false positive results secondary to poor hepatic synthetic function. Less than 30% of the study population had appropriate follow-up investigations for a positive ceruloplasmin result such as ophthalmic examination for KF rings or liver biopsy for copper staining. Ceruloplasmin measurements were requested by a variety of subspecialties, the most common being gastroenterology, neurology and internists. The most common reason for testing was investigation of unexplained liver elevation or established liver disease. Physicians who were not gastroenterologists or neurologists were less likely to perform confirmatory investigations (p<0.01). This suggests lack of familiarity from other subspecialties with the diagnostic algorithm of investigations necessary to confirm or exclude WD.

Serum ceruloplasmin measurement with a cut-off level of less than 0.2g/L had a low pre-test probability of the diagnosis in an unscreened population, with a high prevalence of liver disease with known etiology. Lowering the reference range to 0.14 g/L significantly improved test performance, with a change in sensitivity as all 3 WD cases had a serum ceruloplasmin <0.05g/L. There was a statistically significant relationship between gender, age, and likelihood of a positive ceruloplasmin test result. The mean age of those with a positive level was 36.5 years compared to 45.8 years in those without (p=0.03). 10% of the tested male population had a level low versus 5.5% of the female population (p=0.01). There was a three-fold increase in frequency of ceruloplasmin testing per year over the 7 years studied comparing the tests done in 2003-04 (109) to 2008-2009 (362).

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