Review Article

Mild Hypertransaminasemia in Primary Care

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The liver enzymes, alanine transaminase (ALT) or aspartate transaminase (AST), are commonly used in clinical practice as screening as well as diagnostic tests for liver diseases. ALT is more specific for liver injury than AST and has been shown to be a good predictor of liver related and all-cause mortality. Asymptomatic mild hypertransaminasemia (i.e., less than five times normal) is a common finding in primary care and this could be attributed to serious underlying condition or has transient and benign cause. Unfortunately, there are no good literatures available on the cost-effectiveness of evaluating patients with asymptomatic mild hypertransaminasemia. However, if the history and physical examination do not suggest a clear cause, a stepwise approach should be initiated based on pretest probability of the underlying liver disease. Nonalcoholic fatty liver disease is becoming the most common cause of mild hypertransaminasemia worldwide. Other causes include alcohol abuse, medications, and hepatitis B and C. Less common causes include hemochromatosis, α1-antitrypsin deficiency, autoimmune hepatitis, and Wilson’s disease. Nonhepatic causes such as celiac disease, thyroid, and muscle disorders should be considered in the differential diagnosis. Referral to a specialist and a possible liver biopsy should be considered if persistent hypertransaminasemia for six months or more of unclear etiology.

1. Introduction

The term “liver function tests (LFT)” is commonly used in clinical practice when referring to liver enzymes. However this term is misleading since these tests primarily assess liver injury rather than liver function. In addition, this term also implies that these tests are solely of hepatic origin, but in fact, they are not always specific for the liver. The serum aminotransaminases, alanine transaminase (ALT, formally serum glutamic-pyruvic transaminase (SGPT)) and aspartate transaminase (AST, formally serum glutamic-oxaloacetic transaminase (SGOT)), are frequently obtained in primary care for screening and diagnosis of liver diseases and have therefore led to increased number of asymptomatic patients with mild hypertransaminasemia (i.e., less than five times the upper limit of normal) [1]. A population-based survey in the United States found that hypertransaminasemia was present in 8.9 percent of the population [2]. Although the majority of these individuals will have benign conditions, a subgroup will have underlying significant liver diseases that require further evaluation and therapeutic intervention.

An audit of primary care practices found that these abnormalities are often not adequately investigated, missing an important chance of identifying treatable chronic liver disease [3]. Given the importance and frequency of this issue, the primary care physician should develop a rational approach to the management of mild hypertransaminasemia. There are several published guidelines for the evaluation of asymptomatic hypertransaminasemia [1, 4, 5], which are mainly based on expert opinions, and a very limited number of clinical studies. Understanding the basic disease processes that cause mildly hypertransaminasemia and the epidemiology of each disease can help guide the clinical evaluation and efficient use of laboratory testing.

2. Literature Review

We searched PubMed using the following keywords: “elevated,” “liver tests,” “liver enzymes,” “transaminase,” and “aminotransferase.” The articles were excluded from our paper if they did not pertain to the topic of hypertransaminasemia, or if they were in a language other than English.
The rest of our recommendations are based on data from few retrospective studies, case series (Level B), or expert consensus guidelines (Level C).

3. Transaminases as Markers of Liver Disease

The sera of ALT and AST are the most reliable and sensitive indicators of hepatocellular injury. Their levels can be elevated in a variety of hepatic disorders. ALT, which is present mainly in the cytosol of liver and much less in the other tissues, is the most specific for liver disease [6], whereas AST, which has cytosolic and mitochondrial forms, is less specific for liver disease as it is found, in addition to the liver, in the heart, skeletal muscle, kidneys, brain, pancreas, lungs, leukocytes, and erythrocytes. Although levels of ALT and AST can be extremely elevated (greater than 15 times the upper limit of normal (ULN) in cases of acute liver injury related to drugs, toxins, ischemia, and hepatitis), elevations less than 5 times the ULN are much more common in primary care practice.

Although one study suggested that the majority of asymptomatic individuals (88%) with mild ALT elevations do not have identifiable causes [7], a Scandinavian study of 151 consecutive patients with mild to moderate elevations (42–300 IU/L) of ALT levels for at least 6 months revealed that identifiable causes of liver disease were more common [8]. Diagnoses included NAFLD in 42%, chronic HCV in 15.3%, presumed alcoholic liver disease in 8%; autoimmune hepatitis, primary biliary cirrhosis, and alpha1 antitrypsin deficiency were much less common. Both of these studies were conducted in the era of less sensitive HCV testing and therefore the true prevalence of HCV infection could not be ascertained.

The AST/ALT ratio can suggest certain disease patterns. In most of the liver diseases (acute or chronic), the ratio is less than or equal to 1 such as nonalcoholic fatty liver disease (NAFLD) and viral hepatitis (B or C) [9], whereas, an AST/ALT ratio greater than 2 characteristically is present in alcoholic hepatitis [10] and a ratio greater than 4 suggests Wilson’s disease [11]. In addition, smaller increases in the ratio to values greater than 1.0 suggest the presence of cirrhosis in different causes of nonalcoholic liver diseases such as NAFLD, hepatitis B, and hepatitis C [9]. Therefore, a ratio greater than 1.0 in a cirrhotic patient is not necessarily indicative of alcoholic liver disease. Furthermore, while these ratios are suggestive of certain conditions, there is too much overlap between groups to rely on them exclusively when making a diagnosis.

In addition, aminotransferase levels may be normal in a subset of liver disease despite advanced histologic features. In other words there is a lack of correlation between the level of transaminases and severity of the histologic damage in patients with chronic hepatitis such as nonalcoholic steato-hepatitis [12] and hepatitis C [13, 14].

4. ALT: What Is Normal Value?

ALT is commonly used in clinical practice as a screening, diagnostic, and monitoring test for liver diseases [15]. However, accepted definitions and uniform measurements of the normal range have been an unsettled issue. Therefore, ALT level, which differentiates between asymptomatic persons who have liver disease and those who do not, is still uncertain. Several studies have addressed the ULN of ALT. Two recent reports from United States showed a wide variation in ALT ULN values across laboratories [16, 17]. Some of this variation may be attributed to the use of different instruments or methods to perform the measurement or to differences in reference populations used to define the ULN [18].

The reference ranges for routine laboratory tests are determined based on 2 methods, either on values obtained from healthy individuals or on health outcomes [19]. The former, which is the commonly used method, involves selecting apparently healthy individuals and setting the reference limits (upper and lower ends of the “normal range”) to include the central 95% of values for the test, whereas, the use of health outcome-based reference ranges requires that there is a high degree of standardization across different labs, and a precise relationship exists between adverse health outcome and a discrete level of the lab value.

When using the central 95% of ALT values of the “normal” population, 5% will have abnormal ALT (2.5% are above and 2.5% are below the normal range) which means that 2.5% of normal individual will have abnormally high results. In addition, the current ULN for ALT (40 IU/L in men, 30 IU/L in women) has been established in 1980s when it was introduced as a surrogate marker for screening of hepatitis C among blood donors and before the implementation of anti-HCV testing and restrictive behavioral criteria for donor selection. This explains why ALT ULN is different between labs as those individuals were only “apparently healthy,” and have other reasons for increased ALT. The most important of these are Hepatitis C and NAFLD. Other factors also known to cause increased ALT include medications, herbal supplements, and excessive ethanol ingestion [19]. In a study of blood donors conducted in Milan, Italy, which included individuals at lowest risk of liver disease (no hepatitis B or C, not overweight, alcohol abuse or taking any medications, and those with normal levels of cholesterol, triglycerides, or glucose), the ULN was 30 IU/L in men and 19 IU/L in women [20]. In addition, a similar result for men was obtained in a South Korean outcome study of more than 90,000 men followed for liver related mortality over 8 years, where cut-off value of 30 IU/L was the best predictive of liver disease in men [21]. This new ULN was shown to increase sensitivity for detection of hepatitis C virus among blood donors from 40% to 61% with very slight decline in specificity from 97.6% to 95.5% [20]. Using the old ULN of ALT, 50% of the individuals with increased ALT had no obvious cause for liver injury, but most of them (85%) had evidence of steatosis on liver ultrasound. Therefore, it is clear that the current different ULN used by different labs is insensitive to the presence of liver disease and does not detect limits at which risk of liver mortality seems to increase. In addition, this wide variation in levels considered “abnormal” between different labs leads to differences in the recognition of liver diseases and decision to treat hepatitis B virus [22]. In the other hand, using
the new lower ULN might lead to labeling a large number
of the populations as having liver injury when the cost of
evaluation would be high and its utility is still uncertain.

Several strategies should be used in order to establish the
“healthy” range of ALT of the target population including
using similar lab methods and excluding individuals with
risk factors for liver diseases. We, the authors of this review,
commonly use 30 IU/L for men and 19 IU/L for women as a
reference for ULN of ALT when dealing with patients at high
risk for liver diseases.

5. ALT as Predictor of Health Based Outcomes

While ALT is useful as an initial test in detecting liver disease,
emerging data highlight its potential value as a measure of
overall health and survival. There are several observational
studies which have shown a strong relationship between
ALT activity and liver and all-cause mortality. The strongest
is a population-based study from Korea which included a
cohort of 142,055 participants of ages between 35 and 59
years followed up to 10 years, when death certificates were
used to determine survival and causes of death [21]. In this
study, the risk of death from liver disease started to increase
at ALT value above 20 IU/L. Moreover, increased ALT has
been associated with the risk of death from all-cause or
cardiovascular disease with risk beginning to increase even
at level below current ULN [21, 23, 24]. While mortality
risk may be due to unknown liver disease, it may partly be
explained by the presence of metabolic syndrome in patients
with NAFLD, in addition to alcohol consumption, which are
linked to nonliver health risks.

6. Initial Approach to the Evaluation of
Mild Hypertransaminasemia

The clinical significance of mild hypertransaminasemia must
be interpreted in the context of the clinical presentation. In
general, symptomatic patients (i.e. with signs or symptoms of
chronic liver disease or evidence of hepatic decompensation
e.g. ascites, encephalopathy, coagulopathy, or portal hyper-
tension) should be evaluated and treated in a more expedi-
tious manner than asymptomatic patients with normal phys-
ical exam. Unfortunately, there are no data available on the
cost-effectiveness of evaluating patients with asymptomatic
mild hypertransaminasemia, nor on the natural history of
the potential liver disease in these patients. However, given
the high prevalence of this abnormality in the primary care
settings and the significant costs of an extensive evaluation,
rational stepwise approach should be guided by the pretest
probability of the underlying liver disease, the pattern of
abnormalities, and suggestive features obtained from the
history and physical examination. The following sections will
provide this approach based on the published guidelines
[1, 4, 5] (Figure 1).

A detailed history and physical examination are essential for
the initial evaluation to determine whether the liver
injury is acute or chronic (defined as ≥6 months), the
underlying cause, and associated comorbidities. Important
considerations include:

(i) the presence of any accompanying symptoms such as
abdominal pain, fever, and weight loss or symptoms of
liver dysfunction such as jaundice, confusion, ascites,
and leg swelling;

(ii) the exposure to any medication including prescrip-
tion and over-the-counter medications as well as
herbal therapies;

(iii) occupational exposure to other hepatotoxins and
alcohol consumption;

(iv) risk factors for viral hepatitis including possible par-
tenteral exposures including transfusions, intravenous
and intranasal drug use, tattoos, and sexual activity;

(v) family history of liver disease or other autoimmune
disorders.

The physical examination should include assessment for
signs of metabolic syndrome, a risk factor for NAFLD,
such as hypertension, obesity (body mass index, BMI, and
waist circumference), dyslipidemia (corneal urchus and xan-
thomas), and insulin resistance (acanthosis nigricans). In
addition, patients should be examined for findings suggesting
the presence of liver disease, including the assessment for
stigmata of chronic liver disease (e.g., spider nevi, palmar
erythema, gynecomastia, etc.), splenomegaly (suggestive of
portal hypertension), and ascites.

If the history and physical examination suggest a par-
ticular diagnosis, a targeted testing should follow. On the
other hand, more than 30% of individuals with initially mild
hypertransaminasemia will have normal levels on retesting
[25]. Therefore, if ALT elevation is less than 2 times normal
and if the history and physical examination do not suggest
an etiology or a laboratory error is strongly suspected, it is
reasonable to repeat the test in four weeks [1, 4]. However,
it should be cautioned that hepatitis C may present with
fluctuating elevated liver enzymes levels.

In addition, a period of effective lifestyle changes can be
initiated including complete abstinence from alcohol,
control of diabetes and hyperlipidemia, weight loss in
overweight patients, and stopping or changing potentially
hepatotoxic medications and supplements (Figure 1). Such
lifestyle changes directly impact several of the causes of
mild hypertransaminasemia and may be all that is needed to
correct the abnormalities [11] (Table 1).

6.2. Look for Common Causes. Additional laboratory tests
for common causes of liver injury should be obtained when
history and physical examination show no obvious cause
(Table 1). Hepatitis B, hepatitis C, and hemochromatosis
should be considered [4]. These conditions were found to
be responsible for mild hypertransaminasemia in only 31%
of patients [26]. The majority of the patients (69%) had
unexplained elevations, but there were strong associations
with markers of metabolic syndrome and therefore may rep-
resent NAFLD [26]. This observation was also confirmed in
Mild hypertransaminasemia

Step 1: history and physical examination consider discontinuing hepatotoxic medications and lifestyle changes

The history and physical examination do not suggest a diagnosis (or a laboratory error is not suspected)

The history and physical examination suggest a diagnosis

Step 2: consider common causes (Table 1) and get complete blood counts with platelets, albumin, PT, and INR

Evidence of chronic liver disease or hepatic decompensation

Expeditied evaluation and referral to specialist

Negative initial workup without hepatic decompensation

Consider control of metabolic syndrome including weight loss, exercise, diet, diabetes, and dyslipidemia control

6-month observation with repeat transaminase levels

Normal

Persisting or worsening

Observe

Persistent or worsening

Step 3: consider nonhepatic and rare liver causes (Table 1)

If positive or persistent

Step 4: consider referral to specialist and need for liver biopsy

Targeted evaluation and treatment

Positive initial workup

Consider further testing based on the suspected diagnosis (see Table 1) and referral to specialist

Figure 1: Management algorithm of mild hypertransaminasemia [1, 4, 5].

A previous study showed that NAFLD was responsible of at least 80% of asymptomatic hypertransaminasemia after other causes were ruled out [27]. The measurement of a complete blood count with platelet count, coagulation profile, and albumin should be considered if liver dysfunction is suspected.

6.3. Consider Nonhepatic Causes and Rare Liver Conditions. If the etiology was not reached despite the above workups, observation with lifestyle changes should be undertaken for up to 6 months [4]. If hypertransaminasemia persists or worsens, the patient should be reevaluated and further diagnostic testing should be performed, if necessary. The next step should include testing for nonhepatic causes, including muscles and thyroid diseases, as well as rare causes like celiac disease, based on the clinical scenario (Table 1). In addition, rare liver diseases, including Wilson's disease, Alpha-1 antitrypsin deficiency, and autoimmune hepatitis, should be considered at this stage (Table 1).
Table 1: Causes of mild hypertransaminasemia, clinical clues and initial diagnostic testing [1, 4, 5].

<table>
<thead>
<tr>
<th>Etiology</th>
<th>Clinical clues</th>
<th>Initial diagnostic testing</th>
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<tbody>
<tr>
<td><strong>Common causes</strong></td>
<td></td>
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<tr>
<td>Drugs (including prescribed, over the</td>
<td>(i) Lack of illness prior to taking the drug</td>
<td>History</td>
</tr>
<tr>
<td>counter, illicit drug use and herbals</td>
<td>(ii) Clinical illness or biochemical abnormalities developing after beginning</td>
<td>Accurate history, CAGE questionnaire, AST/ALT ratio, γ-glutamyl</td>
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<td></td>
<td>the drug</td>
<td>transpeptidase level (GGT)</td>
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<td></td>
<td>(iii) Improvement after the drug is withdrawn</td>
<td></td>
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<tr>
<td>Alcohol abuse</td>
<td>Excessive alcohol consumption, AST/ALT ratio ≥ 2.0</td>
<td>Hepatitis B surface antigen (HBsAg), Hepatitis B surface antibody</td>
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<td></td>
<td></td>
<td>(HBsAb), Hepatitis B core antibody (HBCAb)</td>
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<tr>
<td>Nonalcoholic fatty liver disease</td>
<td>Evidence of metabolic syndrome (dyslipidemia, hypertension, diabetes or central</td>
<td>Fasting lipid profile, glucose level; consider ultrasonography to</td>
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<td>(NAFLD)</td>
<td>obesity) (AST/ALT ratio &lt; 1.0)</td>
<td>detect hepatic steatosis</td>
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<td></td>
<td>High risk factors including (Immigration from endemic countries, high risk</td>
<td>Hepatitis C virus antibody testing</td>
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<td></td>
<td>sex behavior, and intravenous drug use)</td>
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<tr>
<td></td>
<td>Parenteral exposure (blood transfusions, intravenous drug use, occupational),</td>
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<td>tattoos, body piercing, and high risk sexual behavior</td>
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<tr>
<td>Hereditary Hemochromatosis</td>
<td>Family history</td>
<td>Transferrin saturation and ferritin levels</td>
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<tr>
<td><strong>Less common causes</strong></td>
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<tr>
<td>Autoimmune hepatitis</td>
<td>Personal of family history of other autoimmune diseases</td>
<td>Immunoglobulin G levels, Serum protein electrophoresis (SPEP),</td>
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<td></td>
<td></td>
<td>antinuclear antibodies (ANA), smooth muscle antibodies (SMA) and</td>
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<td>liver-kidney microsomal antibodies testing (LKMA)</td>
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<tr>
<td>Wilson's disease</td>
<td>Younger than 40 years, neuropsychiatric symptoms, Kayser-Fleischer rings</td>
<td>Serum ceruloplasmin level and ophthalmologist consultation to</td>
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<tr>
<td></td>
<td></td>
<td>rule out Kayser-Fleischer rings</td>
</tr>
<tr>
<td>α1-antitrypsin deficiency</td>
<td>Early-onset emphysema, family history</td>
<td>Serum α1-antitrypsin level and SPEP</td>
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<tr>
<td><strong>Non-Hepatic causes</strong></td>
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<tr>
<td>Muscle disorders</td>
<td>Muscle weakness and pain, strenuous exercise</td>
<td>Creatine kinase (CK) and aldolase levels</td>
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<tr>
<td>Thyroid disorders</td>
<td>Signs and symptoms of hypo- or hyperthyroidism</td>
<td>Thyroid-stimulating hormone (TSH) level</td>
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<tr>
<td>Celiac disease</td>
<td>Diarrhea, abdominal pain, malabsorption</td>
<td>Tissue transglutaminase antibody (TTG) testing</td>
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</table>

6.4. *Referral to a Specialist*. Consultation with a gastroenterologist or hepatologist should be considered for the following groups of patients [5]:

(i) patients with unexplained hypertransaminasemia on two occasions, a minimum of 6 months apart despite lifestyle changes;

(ii) symptoms or signs of liver decompensation (stigmata of chronic liver diseases, ascites, and hepatic encephalopathy);

(iii) evidence of liver dysfunction (high bilirubin, low albumin, and prolonged PT or INR);

(iv) evidence of liver disease where treatment beyond the withdrawal of implicated agent (alcohol or drugs) is warranted, for example, hepatitis B, hepatitis C, autoimmune hepatitis, Wilson's disease, hemochromatosis, and NAFLD.

7. *Indications for Liver Biopsy*

In general, liver biopsy has three major roles: to make a firm diagnosis (or exclude a diagnosis of any serious or significant liver disease), for assessment of prognosis (disease staging), and/or to assist in making therapeutic management decisions. In cases of mild hypertransaminasemia, it is often considered in patients in whom all noninvasive tests were negative or in patients in whom a specific liver disease has been considered but has not yet been confirmed, for example, Wilson's disease and Alpha-1 antitrypsin deficiency. While it remains less likely that the biopsy will provide a diagnosis or lead to changes in management, it is often reassuring for the patient and clinician to know that there is no serious disorder.

8. *Conclusion*

Mild hypertransaminasemia is a common finding in primary care practice. Unfortunately, there are no data available on the cost-effectiveness of evaluating such patients. If the history
and physical examination do not suggest a cause, a stepwise approach should be initiated based on the pretest probability of the underlying liver disease. Patients with an abnormal albumin or prothrombin time or with evidence of chronic liver disease and/or hepatic decompensation should typically have more expeditious evaluations preferably by a specialist. Referral to a specialist is also recommended if unexplained asymptomatic hypertransaminasemia remains elevated for six months or more.

Disclosure

The authors have no relevant financial affiliations to disclose.

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