Esterified Anabolic Androgen-Induced Liver Injury in a Hepatitis C Virus-Positive Patient: A Case Report

Kyrillus S. Shohdy,¹ Rasmia M. El Gohary.²

Abstract

Background: Cases of drug induced liver injury still perplex gastroenterologists due to its wide range of presentations that mimic acute and chronic liver conditions. Moreover, matters get complicated when clinicians face the possibility of drug-induced injury in the presence of pre-existing chronic liver disease. Case: A 69 year-old male who was recently discovered to have a hepatitis C viral infection presented with acute manifestations (mixed cholangio-hepatocellular injury) not fully explained by the underlying chronic disease, we suspected an idiosyncratic reaction from an esterified anabolic androgen. His manifestations have appeared acutely after the drug intake and include acute onset of jaundice, abdominal pain, pruritus and choluria. He was improving on drug discontinuity and conservative measures during his brief hospital stay. Conclusion: The underlying chronic disease constitutes a dilemma in diagnosis of superimposed drug-induced liver injury, as the proof of causality is a daunting task. In such cases, it is tempting to link such new emerging manifestations to be a flare-up of the underlying chronic disease rather than to the drug. However, certain clues helped to point this clinical presentation towards a drug-induced liver injury.

Introduction

Cases of drug induced liver injury (DILI) still perplex gastroenterologists due to its wide range of presentations that mimic acute and chronic liver conditions. The liver is the most common organ prone to toxicity. However, the absence of internationally accepted diagnostic criteria for DILI along with underreporting and underrecognition make the overall incidence variable which ranges from 1 in 10,000 to 1 in 100,000 patients. For instance, in one population-based cohort study, the incidence was 14 cases in 100,000.² It is worth noting that hepatotoxicity was found to be the most common cause behind drug withdrawal or drug usage modification.² The naturally occurring testosterone hormone can be modified for therapeutic use by one of two ways: esterification or alkylations. The alkylated forms can be taken orally and are considered a common cause of liver injury, whereas the esterified forms such as testosterone enanthate (TE) are considered debatable cause for DILI as these drugs are taken illicitly and case reports are the only available evidence.² This case may open

Key Points:
- The clinical and biochemical presentation of drug-induced liver injury mimics a broad array of liver diseases.
- In some instances, liver biopsy can provide unequivocal differentiation between flare-ups of chronic liver diseases and superimposed liver damage caused by medications.
- Short timespan between drug intake and the onset of liver injury manifestations, the pattern of hepatotoxicity and presence of conjugated hyperbilirubinemia could point to the diagnosis of drug-induced liver injury.
- Liver function tests should be performed before starting anabolic steroids, either oral or injectable, in endemic areas of asymptomatic chronic liver diseases.

Figure 1. Structure of testosterone (A) and its esterified form, testosterone enanthate (B)

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Keywords: Drug-Induced Liver Injury; Hepatitis; Steroids; Biological Markers (Source: MeSH, NLM).
Tadalafil not known to be hepatotoxic.21

Table 1. Case Summary.

<table>
<thead>
<tr>
<th>Criteria</th>
<th>Possible Scores</th>
<th>Present Case Score</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Duration of culprit drug therapy</td>
<td>7 days</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time to onset of symptoms</td>
<td>8 days</td>
<td></td>
<td></td>
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<tr>
<td>Clinical presentation</td>
<td>Jaundice, abdominal pain, pruritus, cholina</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time to admission</td>
<td>21 days</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Initial R ratio</td>
<td>3.3 (Mixed pattern)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time to first detection of lab abnormalities</td>
<td>21 days</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Liver function tests at time of admission</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT= 6.9 times ULN*, AST= 5.9 times ULN, ALK P=2.25 times ULN, total bilirubin 30.0 mg/dL, direct 28.0 mg/dL.</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>After 5 days of admission</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ALT, AST and Alk P showed mild decline, total bilirubin 27.0 mg/dL, direct 25.0 mg/dL.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Time to resolution</td>
<td>Not applicable</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Legend: ULN= the upper limit of normal.

The Case

A 69 year-old Egyptian male patient presented to our emergency room with an acute onset of jaundice, abdominal swelling, itching and dark colored urine and pale stools. After 5 days of admission, the total and direct bilirubin were 27.5 mg/dL.

Abdominal examination revealed the presence of generalized abdominal distention, epigastric tenderness, enlargement of the liver with a sharp border and shifting dullness revealed ascites. His past medical history is unremarkable for previous surgeries or chronic morbidities except for an attack of myocardial infarction. His liver function tests (LFTs) showed a mixed hepatitis/cholestasis pattern where alanine transaminase (ALT) was 240 U/L [6.9 times the upper limit of normal], aspartate transaminase (AST) 264 U/L [5.9 times the upper limit of normal], alkaline phosphatase (Alk P) 270 U/L [2.25 times the upper limit of normal], hence the calculated R value is 3.3, total bilirubin 30.0 mg/dL, (direct 28.0 mg/dL), albumin 3.7 g/dL and eosinophils 2%. The abdominal ultrasound revealed hepatomegaly with no dilatation of the intrahepatic biliary tract radicles with normal sized kidneys and his autoimmune and virology screening was negative except that his HCV RNA was qualitatively positive. The patient was not aware of being HCV positive before. We restricted our screening to the recommendations of the American College of Gastroenterology clinical guidelines for investigating a case of suspected DILI that included anti-nuclear antibody, anti-smooth muscle antibody, IgG level, Anti-HAV IgM, HBsAg, anti-HBc IgM, anti-HCV and HCV RNA.9 On follow-up, after 5 days of admission, the total and direct bilirubin were 27.5 mg/dL.

Table 2. RUCAM criteria to prove causality in DILI cases.

<table>
<thead>
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<th>Possible Scores</th>
<th>Present Case Score</th>
<th>Rationale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Time between drug start and liver enzyme elevations</td>
<td>+1 to +2</td>
<td>+2</td>
<td>Receiving the drug for the first time is given 2 points if the time to onset is 5 to 90 days.</td>
</tr>
<tr>
<td>Course of the reaction</td>
<td>-2 to +5</td>
<td>+1</td>
<td>Mild resolution of LFTs in 20 days after drug discontinuity</td>
</tr>
<tr>
<td>Risk factors</td>
<td>0 to +2</td>
<td>+1</td>
<td>Age ≥ 55 yrs.</td>
</tr>
<tr>
<td>Concomitant drugs</td>
<td>-3 to 0</td>
<td>0</td>
<td>Tadalafil not known to be hepatotoxic.21</td>
</tr>
<tr>
<td>Competing non-drug causes</td>
<td>-5 to +2</td>
<td>-1</td>
<td>Potential HCV activity</td>
</tr>
<tr>
<td>Previous Information on hepatotoxicity of the drug</td>
<td>0 to +2</td>
<td>+2</td>
<td>See discussion section.</td>
</tr>
<tr>
<td>Rechallenge</td>
<td>-2 to +3</td>
<td>0</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Score</td>
<td>8+ points definite</td>
<td>6-8 points probable</td>
<td>3-5 points possible</td>
</tr>
</tbody>
</table>
In addition, the HLA-B*5701 genotype has been linked with flucloxacillin induced liver injury. 20 It is very important to those with chronic liver disease, for example, Abacavir which is an antiviral used in human immunodeficiency virus (HIV) treatment, can cause severe hypersensitivity reactions and hepatotoxicity. These reactions have been found to occur in patients with human leukocyte antigen-B*5701 and -1502 alleles. 21 Genetic screening prior to Abacavir intake is strongly advised. 22,23 Though these genetic linkage studies can predict what was thought to be unpredictable idiosyncrasies, there is still a long way to reach widely accepted predictive biomarkers.

Conclusion

Two clinical notes learnt from this case; first, LFTs are recommended before starting anabolic steroids either oral or injectable in endemic areas with asymptomatic chronic liver diseases such as HCV. Second, in any case of acute liver injury within endemic areas, stigmata of chronic liver disease have to be investigated thoroughly by clinical, biochemical and radiological means. Physicians have to take into consideration that HCV as well as other chronic diseases e.g. alcoholic liver disease and nonalcoholic fatty liver disease (NAFLD) leave the liver susceptible to steroid toxicity. In contrast to alcoholic hepatitis and HBV, HCV patients do not experience icteric flares, which along with the history of hepatotoxic agent intake, latency to DILI onset, and biochemical and histological features at presentation raise the index of suspicion to DILI. However, due to diagnostic limitations it is difficult to distinguish with a high-degree of certainty between a spontaneous disease flare-up and a DILI episode.

mg/dl & 25 mg/dl respectively and LFTs were ALT 186 U/L, AST 203 U/L, Alk P 189 U/L.

The symptoms of the patient were improving slowly on withdrawal of TE. Simultaneously, intravenous fluids and N-acetylcysteine were administered while monitoring for the development of coagulopathy or encephalopathy. There were no clinical benefits found in the literature regarding the use of ursodeoxycholic acid (UDCA) in the mixed hepatotoxicity pattern except for lowering alkaline phosphatase levels. 17 The patient left against medical advice so a follow-up could not be reported. Written informed consent was obtained from the patient for publication of this case report.

Discussion

The clinical and biochemical presentation of drug-induced liver injury mimics a broad array of liver diseases. However, the most common clinical presentation resembles hepatocellular jaundice or cholestatic liver disease. According to the LiverTox®, (Available from: http://livertox.nih.gov/AndrogenicSteroids.html, updated 2015 June 23; cited 2015 July 21) toxicity with injectable anabolic steroids (esterified testosterone) is not well documented. In the review of the literature, we found two cases that developed liver toxicity (acute cholestasis) while using esterified testosterone, 4,15 and one case that developed a hepatic adenoma. 16 Moreover, preclinical studies showed that long term use of esterified testosterone like testosterone enanilate can cause an increase in liver enzymes and a decrease in HDL-cholesterol. 17 On the other hand, one study by Marquardt et al in 1964 showed failure of esterified anabolic steroids (non C17-alkylated) to produce normal liver function tests. 4 Further studies are needed to assess the short and long term effects of esterified testosterones on the liver.

The differential diagnoses of this case includes acute viral hepatitis, autoimmune hepatitis and other less common viruses e.g. cytomegalovirus and Epstein-Barr virus. Acute hepatitis C can masquerade as DILI in 1.3% of cases especially with acute hepatocellular injury and suggestive HCV RNA testing. 20 However, these two findings do not apply to our case, for example, Chalasani et al. recruited 300 cases with suspected DILI in a prospective study and found that DILI is an unlikely diagnosis with only 9 positive cases (3%) and in 4 of them the final diagnosis turned out to be an HCV infection. 18 Another study, recruited 570 cases, has excluded 59 cases for an alternative cause of injury which was viral hepatitis on 11 occasions. A total of 446 (98%) cases were deemed to be idiosyncratic hepatotoxicity. 18 Although viral hepatitis is a valid differential diagnosis in suspected DILI cases, studies have shown that it is a less likely cause of injury in idiosyncratic liver injury.

The existence of underlying liver disease can still hinder a straightforward diagnosis of DILI in general, since the flare-up of the chronic disease caused by viral hepatitis cannot be ruled out completely. In some instances, liver biopsy can provide unequivocal differentiation between flare-ups and superimposed damage by the offending drug. Other important clues are the short time period between the culprit drug intake and the onset of clinical manifestations, the pattern of hepatotoxicity and presence of conjugated hyperbilirubinemia. 16 We depended on these three previous clues for considering DILI as a possibility in this case. We have also implemented the Roussel Uclaf Causality Assessment Method (RUCAM), which was developed in 1993 by a group of international experts; our case achieved a possible (+5) score (refer to Table 2). An interesting review from the Spanish Group for the Study of Drug-Induced Liver Disease points out the strengths and weaknesses of the available causality assessment methods. 19

The prevention of idiosyncratic drug reactions is a daunting task; however the future is promising with progress towards the development of biomarkers, where the genetic profile of every person can be screened to determine whether a certain drug can cause an idiosyncratic toxicity. This advancement will be very important to those with chronic liver disease, for example, Abacavir which is an antiviral used in human immunodeficiency virus (HIV) treatment, can cause severe hypersensitivity reactions and hepatotoxicity. These reactions have been found to occur in patients with human leukocyte antigen-B*5701 and genetic screening prior to Abacavir intake is strongly advised by the Food and Drug and Administration (FDA) to prevent such severe reactions. 20 In addition, the HLA-B*5701 genotype has been linked with fluvoxacinil induced liver injury. 21 Though these genetic linkage studies can predict what was thought to be unpredictable idiosyncrasies, there is still a long way to reach widely accepted predictive biomarkers.
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Case Report

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