Acute Liver Injury: Due to a Rare Case Accompanied by Cardiac and Pulmonary Complications

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Abstract

Background: One of the most important causes of acute hepatitis is vascular pathologies of the liver. There are several well-recognized forms of vascular injury to the liver, including Budd-Chiari syndrome, ischemic hepatitis, hepatic infarction and passive congestion due to hemodynamic instability. We present a case of acute liver injury accompanied by hemodynamic and respiratory pathologies.

Case: A 53-year-old lady was admitted to the emergency department with complaints of abdominal pain, nausea, vomiting for a week. Gastrointestinal examination was normal. Laboratory tests revealed mild thrombocytopenia, prolonged international normalized ratio (INR) and transaminases were ten-fold higher. In abdominal ultrasonography, no signs of chronic liver failure and vascular pathologies. Abdominal computed tomography (CT) revealed no pathology. Transthoracic echocardiography disclosed severe mitral regurgitation and dilatation of both atria and right ventricle. Pro-BNP level was increased. Thorax CT demonstrated filling defect compatible with pulmonary thromboembolism (PTE). The patient was accepted PTE, and enoxaparin treatment was initiated. Under the anticoagulant and diuretic treatments, the patient’s orthopnea disappeared, control blood values declined.

Conclusion: Acute liver injury can be the initial presentation in a patient with severe valvular disease and acute PTE.

Keywords
Cardiac decompensation, Echocardiography, Heart failure, Hepatitis, Severe mitral regurgitation, Pulmonary thromboembolism

Abbreviations
ALP: Alkaline Phosphatase; ALT: Alanine Aminotransferase; AST: Aspartate Aminotransferase; CT: Computed Tomography; CTPA: CT Pulmonary Angiography; ECG: Electrocardiography; GGT: Gamma Glutamyl Transferase; INR: International Normalized Ratio; LDH: Lactate Dehydrogenase; Pro-BNP: Prohormone B-Type Natriuretic Peptide; PTE: Pulmonary Thromboembolism

Introduction

Hepatitis refers to an inflammatory condition of the liver. It’s commonly caused by a viral infection, but there are other possible causes of hepatitis. These include autoimmune hepatitis and hepatitis that occurs as a secondary result of medications, drugs, toxins, and alcohol [1]. Henrion, et al. reported that acute hepatitis may also develop in hemodynamic instability such as congestive heart failure, acute heart failure, chronic respiratory failure and septic/toxic shock [2]. Severe acute liver injury defines a syndrome characterised by markers of liver damage (elevated serum transaminases) and impaired liver function (jaundice and INR > 1.5) which usually precedes clinical encephalopathy [3]. Here, we present a case of acute liver injury accompanied by hemodynamic and respiratory pathologies.

Case Report

A 53-year-old morbidly obese lady was admitted to the emergency department with complaints of abdominal pain, nausea, vomiting, weakness, loss of appetite for a week. Her medical history included hypothyroidism and mitral regurgitation. She didn’t
have a history of smoking or exposure to any kind of chemical and hepatotoxic food. She had a history of drinking two beers a week. She had a temperature of 37 °C, a pulse rate of 115 beats/minute, a respiratory rate of 24 breaths/min, in room air fingertip saturation of 96% and a blood pressure of 115/60 mmHg. In the respiratory system examination, inspiratory crackles were heard in the bilateral lower zones. A 2/6 systolic murmur was heard over the mitral area. Pretibial 2+ edema was present. There was no feature in the gastrointestinal examination. There was atrial fibrillation with high ventricular rate in her ECG. Cardiothoracic index was found to be increased in posteroanterior chest X-ray. Hemogram examination showed mild microcytic anemia (hemoglobin: 11 g/dL) and mild thrombocytopenia (platelet count: 109400/μL). In the coagulation panel, international normalized ratio (INR) was prolonged (INR: 2.14), and in the biochemical parameters transaminases were ten-fold higher (AST: 695 U/L, ALT: 983 U/L). Serum cholestatic markers are elevated (ALP: 174 U/L, GGT: 326 U/L, LDH: 717 U/L). Serum bilirubin levels slightly high. The patient with acute hepatitis didn’t consume specific medication and food. Serum viral serologies were negative. Serum transaminase levels were found to be normal in the blood tests performed during the cardiology examination two months ago. In abdominal ultrasonography, no signs of chronic liver failure and vascular pathologies. Liver sizes slightly increased and edematous appearance in perihepatic area. Hepatic veins were normal diameters. Inferior vena cava dilated (IVC > 2 cm) and when the respiratory collapse was less than 50%. Abdominal computed tomography (CT) showed no significant pathology. Autoimmune serologic tests (antinuclear antibody, liver kidney microsomal autoantibody type-1, anti-mitochondrial antibody, anti-smooth muscle antibody) were normal. Intravenous furosemide infusion was initiated due to congestive heart failure, and intravenous atrioventricular nodal blocking agents were applied to treat high ventricular rate. Transthoracic and transoesophageal echocardiogram disclosed normal left ventricular systolic function, severe mitral regurgitation, dilatation of both atria and right ventricle, severe pulmonary hypertension (estimated systolic pulmonary artery pressure was 59 mmHg). And also Pro-BNP level was increased (Pro-BNP: 4281 pg/mL). The patient’s current hepatitis table was evaluated in favor of congestive hepatopathy secondary to cardiac decompensation. For the aetiology of cardiac decompensation, thorax CT pulmonary angiography (CTPA) was performed. In the CTPA study, filling defect compatible with pulmonary thromboembolism was observed in lower lobe segment branch of right pulmonary artery. There was also pleural effusion in right hemithorax and typical pulmonary infarction (Figure 1). The patient was diagnosed with having acute submassive pulmonary thromboembolism, and enoxaparin treatment was initiated. Under the anticoagulant, diuretic and symptomatic treatments, the patient’s orthopnea disappeared, control transaminases regressed (AST:85 U/L, ALT:223 U/L), INR returned to normal. Thrombocytopenia improved (Table 1).

**Discussion**

This case illustrates how the combination of pulmonary embolism and severe valvular heart disease can present as acute liver injury. The circulatory system is a complex interaction between multiple organs. Circulatory dysfunction occurs when one or more organs begin to fail and disrupt adequate delivery of blood to the body [4]. A pathology that occurs in the circulatory system directly affects other organs such as the liver. The majority of cases of ischemic hepatitis occur in the setting of congestive heart fail-

![Figure 1](image-url)
ogy such as Budd-Chiari syndrome, hepatic sinusoidal obstruction syndrome etc. With signs of congestive heart failure (orthopea, in abdominal ultrasound; liver sizes slightly increased and edematous appearance in perihepatic area, inferior vena cava dilated (IVC > 2 cm), when the respiratory collapse was less than 50%) and increased pro-BNP level in mind, trans-thoracic and transoesophageal echocardiography revealed severe eccentric mitral regurgitation (Figure 2) due to rheumatic mitral valve disease, severe pulmonary arterial hypertension. Despite adequate diuresis in response to diuretics and AV nodal blocking agents, the patient’s orthopnoea persisted. Congestive hepatopathy refers to hepatic manifestations attributable to passive hepatic congestion, as occurs in patients right-sided heart failure. Passive congestion often coexists with reduced cardiac output, making their relative contributions to hepatic injury interwined [7]. We hypothesise that the pulmonary thromboembolism was the last hit that resulted in acute liver failure.

The hypotension is the most important clinical finding of cardiovascular collapse caused by massive pulmonary embolism. Early mortality was significantly higher in patients with right ventricular dysfunction than echocardiography despite hypotension [8]. Aslan, et al. was suggested that there may be a re-

Table 1: In blood chart.

<table>
<thead>
<tr>
<th></th>
<th>1st day</th>
<th>4th day</th>
<th>10th day</th>
</tr>
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<tr>
<td>Platelet (count/μL)</td>
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<td>96300</td>
<td>144000</td>
</tr>
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<tr>
<td>Aspartate aminotransferase (U/L)</td>
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<tr>
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<td>983</td>
<td>223</td>
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<tr>
<td>Lactate dehydrogenase (U/L)</td>
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<td>291</td>
</tr>
<tr>
<td>Gamma glutamyl transferase (U/L)</td>
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<td>326</td>
<td>155</td>
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<tr>
<td>Alkaline phosphatase (U/L)</td>
<td>127</td>
<td>174</td>
<td>94</td>
</tr>
</tbody>
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Figure 2: Transoesophageal echocardiogram in apical 4 chamber view depicting severe eccentric mitral regurgitation (white arrow). LA: left atrium; LV: left ventricle.
There is no conflict of interest in connection with this paper.

All of the authors declare that they have all participated in the design, execution, and analysis of the paper, and that they have approved the final version.

References