WHAT'S YOUR DIAGNOSIS?

Gastrointestinal Symptoms and Lactic Acidosis in a Chronic Marijuana User

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A patient presented with diffuse abdominal pain and a history of frequent cannabis use, a diet lacking in meat and fish, and an increase in consumption of simple carbohydrates in the past year.

A 57-year-old woman with a history of traumatic brain injury, posttraumatic stress disorder, depression, migraines, hypothyroidism, and a hiatal hernia repair presented to the emergency department with a 1-day history of nausea, vomiting, and diffuse abdominal pain. She reported that her symptoms were relieved by hot showers. She also reported having similar symptoms and a previous gastric-emptying study that showed a slow-emptying stomach. Her history also consisted of frequent cannabis use for mood and appetite stimulation along with eliminating meat and fish from her diet, an increase in consumption of simple carbohydrates in the past year, and no alcohol use. Her medications included topiramate 100 mg and clonidine 0.3 mg nightly for migraines; levothyroxine 200 mcg daily for hypothyroidism; tizanidine 4 mg twice a day for muscle spasm; famotidine 40 mg twice a day as needed for gastric reflux; and bupropion 50 mg daily, citalopram 20 mg daily, and lamotrigine 25 mg nightly for mood.

The patient's physical examination was notable for bradycardia (43 beats/min) and epigastric tenderness. Admission laboratory results were notable for an elevated lactic acid level of 4.8 (normal range, 0.50-2.20) mmol/L and a leukocytosis count of 10.8×10⁹ cells/L. Serum alcohol level and blood cultures were negative. Liver function test, hemoglobin A₁c, and lipase test were unremarkable. Her electrocardiogram showed an unchanged right bundle branch block. Chest X-ray, computed tomography (CT) of her abdomen/pelvis and echocardiogram were unremarkable.

What is your diagnosis?

How would you treat this patient?

This patient was diagnosed with gastrointestinal beriberi. Because of her dietary changes, lactic acidosis, and bradycardia, thiamine deficiency was suspected after ruling out other possibilities on the differential diagnosis (Table). The patient's symptoms resolved after administration of high-dose IV thiamine 500 mg 3 times daily for 4 days. Her white blood cell count and lactic acid level normalized. Unfortunately, thiamine levels were not obtained for the patient before treatment was initiated. After administration of IV thiamine, her plasma thiamine level was > 1,200 (normal range, 8-30) nmol/L.

Her differential diagnosis included infectious etiology. Given her leukocytosis and lactic acidosis, vancomycin and piperacillin/tazobactam were started on admission. One day later, her leukocytosis count doubled to 20.7×10⁹ cells/L. However, after 48 hours of negative blood cultures, antibiotics were discontinued.

Small bowel obstruction was suspected due to the patient's history of abdominal surgery but was ruled out with CT imaging. Similarly, pancreatitis was ruled out based on negative CT imaging and the patient's normal lipase level. Gastroparesis also was considered because of the patient's history of hypothyroidism, tobacco use, and her prior gastric-emptying study. The patient was treated for gastroparesis with a course of metoclopramide and erythromycin without improvement in symptoms. Additionally, gastroparesis would not explain the patient's leukocytosis.

Cannabinoid hyperemesis syndrome (CHS) was suspected because the patient's symptoms improved with cannabis use. The patient was placed on a high-fat low-carbohydrate diet and her symptoms improved. She was also treated with a course of metoclopramide and erythromycin without improvement in symptoms. Additionally, gastroparesis would not explain the patient's leukocytosis.
discontinuation and hot showers.\textsuperscript{1} In chronic users, however, tetrahydrocannabinol levels have a half-life of 5 to 13 days.\textsuperscript{2} Although lactic acidosis and leukocytosis have been previously reported with cannabis use, it is unlikely that the patient would have such significant improvement within the first 4 days after discontinuation.\textsuperscript{1,3,4} Although the patient had many psychiatric comorbidities with previous hospitalizations describing concern for somatization disorder, her leukocytosis and elevated lactic acid levels were suggestive of an organic rather than a psychiatric etiology of her symptoms.

**DISCUSSION**

Gastrointestinal beriberi has been reported in chronic cannabis users who present with nausea, vomiting, epigastric pain, leukocytosis, and lactic acidosis; all these symptoms rapidly improve after thiamine administration.\textsuperscript{3,6} The patient's dietary change also eliminated her intake of vitamin B\textsubscript{12}, which compounded her condition. Thiamine deficiency produces lactic acidosis by disrupting pyruvate metabolism.\textsuperscript{7} Bradycardia also can be a sign of thiamine deficiency, although the patient's use of clonidine for migraines is a confounder.\textsuperscript{8}

Chronically ill patients are prone to nutritional deficiencies, including deficiencies of thiamine.\textsuperscript{7,9} Many patients with chronic illnesses also use cannabis to ameliorate physical and neuropsychiatric symptoms.\textsuperscript{2} Recent reports suggest cannabis users are prone to gastrointestinal beriberi and Wernicke encephalopathy.\textsuperscript{5,10} Treating gastrointestinal symptoms in these patients can be challenging to diagnose because gastrointestinal beriberi and CHS share many clinical manifestations.

The patient's presentation is likely multifactorial resulting from the combination of gastrointestinal beriberi and CHS. However, thiamine deficiency seems to play the dominant role.

There is no standard treatment regimen for thiamine deficiency with neurologic deficits, and patients only retain about 10 to 15\% of intramuscular (IM) injections of cyanocobalamin.\textsuperscript{11,12} The British Committee for Standards in Haematology recommends IM injections of 1,000 mcg of cyanocobalamin 3 times a week for 2 weeks and then reassess the need for continued treatment.\textsuperscript{13} The British Columbia guidelines also recommend IM injections of 1,000 mcg daily for 1 to 5 days before transitioning to oral repletion.\textsuperscript{14} European Neurology guidelines for the treatment of Wernicke encephalopathy recommend IV cyanocobalamin 200 mg 3 times daily.\textsuperscript{15} Low-level evidence with observational studies informs these decisions and is why there is variation.

The patient's serum lactate and leukocytosis normalized 1 day after the administration of thiamine. Thiamine deficiency classically causes Wernicke encephalopathy and wet

### TABLE Differential Diagnosis for Nausea, Vomiting, and Epigastric Pain

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Characteristics</th>
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<tbody>
<tr>
<td>Cannabinoid hyperemesis syndrome</td>
<td>Symptoms manifest in chronic cannabis users and are relieved by hot water</td>
</tr>
<tr>
<td>Infection</td>
<td>Multiple pathogens; history of exposure, positive blood cultures, and nidus of infection</td>
</tr>
<tr>
<td>Gastrointestinal beriberi</td>
<td>Diagnosis of exclusion, due to thiamine deficiency; leukocytosis and elevated lactic acidosis can be present; symptoms improve after thiamine administration</td>
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<tr>
<td>Gastroparesis</td>
<td>Can be idiopathic or due to another condition like diabetes mellitus or hypothyroidism; diagnosis made by gastric-emptying study</td>
</tr>
<tr>
<td>Pancreatitis</td>
<td>Requires 2 of 3 criteria: epigastric abdominal pain, elevated serum amylase or lipase &gt; 3 times upper limit of normal, or compatible imaging studies</td>
</tr>
<tr>
<td>Psychiatric (eg, rumination syndrome)</td>
<td>Other psychiatric comorbidities are usually present; normal gastric-emptying study</td>
</tr>
<tr>
<td>Small bowel obstruction</td>
<td>History of abdominal surgery; inability to pass flatus or stool; obstruction noted on imaging</td>
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</tbody>
</table>
beriberi. The patient did not present with Wernicke encephalopathy’s triad: ophthalmoplegia, ataxia, or confusion. She also was euvoletic without signs or symptoms of wet beriberi.

CONCLUSIONS
Thiamine deficiency is principally a clinical diagnosis. Thiamine laboratory testing may not be readily available in all medical centers, and confirming a diagnosis of thiamine deficiency should not delay treatment when thiamine deficiency is suspected. This patient’s thiamine levels resulted a week after collection. The administration of thiamine before sampling also can alter the result as it did in this case. Additionally, laboratories may offer whole blood and serum testing. Whole blood testing is more accurate because most bioactive thiamine is found in red blood cells.

Author disclosures
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References