Alpha Fetoprotein-Secreting Gastric Cancer in the Setting of Chronic Hepatitis B: The Role of Endoscopy

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Abstract

Alpha-fetoprotein (AFP) is a common tumor marker for hepatocellular carcinoma (HCC). We report a unique case of an AFP-producing gastric cancer in an elderly man with known chronic hepatitis B (HBV) and a markedly elevated AFP. Thorough diagnostic imaging failed to reveal a HCC, but a gastric adenocarcinoma was found on endoscopy. After successful resection, the serum AFP normalized. On histopathologic examination, the cancer demonstrated focal positive staining for AFP, in keeping with a hepatoid gastric adenocarcinoma. Endoscopy should be considered in clinical situations where increased serum AFP is detected but no HCC is found.

Introduction

Alpha-fetoprotein (AFP) is a common tumor marker for hepatocellular carcinoma (HCC). Rising serum AFP in a patient with cirrhosis or hepatitis B (HBV) should raise suspicion of a developing HCC. However, AFP may be elevated in other malignancies, of which gastric cancer is the most common after HCC and tumors of gonadal origin.1 AFP-producing gastric cancer (AFPGC) has been reported sporadically since 1970,2-6 including several recent case series from Asia.7-9 Our report adds to the growing literature of this rare entity and illustrates the importance of considering endoscopy in a patient with elevated AFP when investigative efforts for HCC are fruitless.

Case Report

An 81-year-old Asian man was followed by his gastroenterologist for chronic HBV and was treated with tenofovir, resulting in undetectable HBV viral load. He did not have cirrhosis or other comorbidities. The patient complained of early satiety, anorexia, weight loss (>4.5 kg), and right-sided chest pain over several months on a follow-up visit. His physical examination showed normal cardiovascular and respiratory function and no appreciable lymphadenopathy. His abdominal exam was benign, with no detectable ascites, no hepatosplenomegaly, and no masses. Laboratory investigations demonstrated a normocytic anemia (hemoglobin 98 g/L, previously 140 g/L) and an elevated AFP of 45 mcg/L (normal: <4.4 mcg/L), while his remaining bloodwork was within normal limits. Given his underlying HBV, exhaustive efforts were directed at identifying a hepatic neoplasm, but computed tomography (CT) and magnetic resonance imaging (MRI) failed to reveal a liver lesion. However, his AFP gradually increased to 540 mcg/L over the ensuing 2 months.

The patient then presented with an acute upper gastrointestinal bleed. He described a 1-week history of melena with no hematemesis or hematochezia. Laboratory tests revealed a hemoglobin of 80 g/L, and normal elec-
Role of Endoscopy in AFP Gastric Cancer

A patient with elevated urea of 9.1 mmol/L was undergoing endoscopy, which revealed a 3-cm ulcerated mass confined to the antrum with no extension into the pylorus (Figure 1). Biopsies showed an invasive, moderately differentiated gastric adenocarcinoma with intestinal differentiation. Larger cells were present, in keeping with a focal hepatoid differentiation (Figure 2). Immunohistochemical staining was negative for Her2/neu and focally positive for AFP.

Staging CT showed no metastatic spread of the disease. The patient received neoadjuvant chemotherapy and subsequent partial gastrectomy. Postoperatively, the patient’s AFP levels normalized, and have remained normal nearly 2 years later.

Discussion

AFP Gastric Cancers are rare, accounting for only 1-6% of gastric cancers. They are often associated with liver and lymph node metastases and a poorer prognosis, and outside Japan and parts of Asia, they are underrecognized. Our patient had chronic HBV, while in previous case series of AFP Gastric Cancers, patients with chronic viral hepatitis, HCC, and/or fatty liver were excluded or their hepatitis statuses were not clearly documented.

Pathologic examination of our patient’s biopsy demonstrated focal hepatoid differentiation consistent with hepatoid adenocarcinoma of the stomach (HAS). This term was originally proposed by Ishikura et al to describe primary gastric cancers with hepatocytic differentiation and large production of AFP. HAS is classified similarly to other AFP Gastric Cancers, but this variant only accounts for 0.38-0.78% of all gastric cancers. HAS predominantly affects males (3.2:1), and the average age of 63.3 years. Given its clinical and pathologic similarities to HCC, prompt diagnosis of hepatoid adenocarcinoma can be difficult, especially in endemic areas with a high incidence of HCC.

As in our case report, previous reports have found that with appropriate treatment of AFP Gastric Cancers such as surgery and/or chemotherapy, AFP levels will decrease. Conversely, with recurrent disease, AFP levels will increase again, suggesting that AFP can serve as a surrogate marker for disease burden. The presence of liver metastases are an important prognostic factor when compared to those with non-AFP Gastric Cancers, suggesting that decreasing AFP levels in AFP Gastric Cancers may be associated with better quality of life and presumably improved survival.

Because of the obvious risk factor of HBV for potentially causing HCC in our case, initial investigative efforts were directed at detecting a HCC. A recently reported case described a patient who also had an unexplained elevated AFP level in the setting of chronic HBV cirrhosis, where variceal screening revealed an AFP Gastric Cancer. In contrast, because our patient had no known cirrhosis, he would not have undergone similar screening.

Although AFP Gastric Cancers have been recognized in Japan and parts of Asia for more than 40 years, there are few publications about AFP Gastric Cancers outside of Asia. We highlight that clinicians should consider this entity in geographic locations with growing Asian immigrant populations. Given the aggressive nature of AFP Gastric Cancers, this malignancy should be included in the differential diagnosis of an elevated AFP. Endoscopy is an appropriate next test if initial imaging is negative.
Disclosures

Author contributions: S. Ip drafted the manuscript. DF Schaeffer provided the pathologic data and photographs. EM Yoshida and WCP Kwan critically revised and approved the manuscript. EM Yoshida is the article guarantor.

Financial disclosure: None to report.

Informed consent was obtained for this case report.

Received: October 28, 2014; Accepted: February 17, 2015

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