Pulmonary hepatoid adenocarcinoma

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Hepatoid adenocarcinoma (HAC) is a rare extrahepatic tumor that morphologically resembles hepatocellular carcinoma. It usually originates in the gastrointestinal tract, but can arise elsewhere. We describe our experience of a case of pulmonary HAC associated with an aggressive course and present available data on HACs utilizing the available literature and the National Cancer Institute Surveillance, Epidemiology, and End Results (SEER) database.

CASE PRESENTATION

Our patient was a 61-year-old man with a 40-pack-year smoking history and severe chronic obstructive pulmonary disease who was referred for evaluation of a suspicious lung nodule. Computed tomography scan revealed a spiculated mass measuring 2.3 cm in the right upper lobe. Percutaneous biopsy revealed pulmonary HAC with tumor cell positivity for thyroid transcription factor 1. An integrated whole-body positron emission tomography–computed tomography scan showed the mass to have a standard uptake value of 4.8. No mediastinal activity was noted. Brain magnetic resonance imaging demonstrated no metastases.

He underwent video-assisted thoracoscopic right upper lobectomy with mediastinal lymph node sampling. Final pathologic review revealed a moderately differentiated, pulmonary HAC (Figure 1). The tumor stained positive for thyroid transcription factor 1, hepatocyte paraffin antigen-1, cytokeratin 7, cytokeratin 8, carcinoembryonic antigen, and cytokeratin 19, and was negative for alpha fetoprotein (AFP), p40, napsin, and cluster of differentiation 10. Final pathological stage was pT1b N0 M0.

Within 3 months after surgery he developed increasing pain in his pelvis, back, and neck. MRI of the spine and pelvis showed diffuse bony disease in multiple areas in the spine and sacrum. A biopsy of the sacral lesion showed metastatic HAC. The patient received radiation therapy to the symptomatic bony metastases but developed progressive weakness and was ultimately transferred to hospice care. He died 6 months after diagnosis.

DISCUSSION

Extrahepatic HACs are described as rare and aggressive tumors most commonly originating in the gastric region (63%). Histologically, these tumors appear as lightly acidophilic cells with abundant cytoplasm and can be diagnosed with fine needle aspiration. HACs follow an aggressive clinical course with a reported overall median survival of 11 months (range, 1-116 months) and a 1-year survival rate of 55%. Primary pulmonary HACs are

Central Message
Pulmonary hepatoid adenocarcinoma is a rare and aggressive tumor with poor overall prognosis. Before surgery, a comprehensive workup is required to ensure no metastatic disease is present.

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extremely rare, with approximately 36 cases reported in the literature. According to previous reports, pulmonary HACs show a predominance in men (92%) and usually present as stage III or stage IV disease. They are mostly seen in heavy smokers. In general, these tumors are quite large at the time of diagnosis, are usually diagnosed at an advanced stage, and are associated with poor outcomes.

We queried the SEER database from 1988 to 2014. Using this database, we identified 41 cases of primary pulmonary HAC (Table 1). Mean age was 63.6 years. Mean tumor size was 8.2 cm. Eighty-two percent of tumors were stage III or IV at presentation. Median overall survival for the entire cohort was 5 months (95% confidence interval, 3.1-6.9 months), with 1- and 3-year survival rates of only 35% and 14%, respectively.

Our report and review of the literature is the largest to date concerning pulmonary HACs. Although previous literature has also suggested that this tumor usually has an aggressive clinical course, prognostic indicators have been difficult to identify. The significance of an elevated AFP level is unknown. The majority of cases have demonstrated elevated serum AFP and/or positive AFP expression on histopathologic analysis. Lack of expression of AFP does not appear to be reliably associated with better outcomes. In our patient, serum AFP levels were not obtained, and the tumor showed no expression of AFP.

There is no standard treatment for HAC. Although advanced disease will usually be treated by platinum-based chemotherapy and/or radiation treatment, patients with localized disease should undergo surgical resection.

It appears that the poor overall survival is a result of the advanced stage at diagnosis for most patients. Mediastinal tissue sampling should be considered before surgery because so many patients present with advanced disease, but there is no available evidence about how frequently this sampling will lead to upstaging. Clinicians should be aware that HACs will tend to have a very aggressive course and a much worse behavior than most non–small cell lung cancers.

CONCLUSIONS
Primary lung HACs are rare tumors with limited data concerning clinical presentation, management, and outcomes. Available literature and a SEER database analysis show that this tumor generally has an extremely poor prognosis. A comprehensive workup and accurate diagnosis is important before any consideration of surgery.

References