Widely Metastatic Parotid Acinic Cell Carcinoma to Bone and Liver: A Case Report, Review of Literature, and Review of Diagnostic Challenges

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Abstract

Distant metastasis of salivary gland neoplasms is a rare occurrence. Generally, high-grade salivary malignancies such as salivary duct carcinoma and high-grade mucoepidermoid carcinoma, as well as tumors located in the submandibular gland, show a higher likelihood of metastasizing. Acinic cell carcinoma is an uncommon salivary gland neoplasm that typically occurs in the parotid gland and is considered a low-grade malignancy. Metastasis is unusual and, if present, predominantly involves the lung and bones. However, vertebral metastasis is exceedingly rare. Here, we describe a rare case of recurrent acinic cell carcinoma of the parotid gland with widespread metastases to multiple bones, including the vertebrae, and to the liver, which was diagnosed in part by cytology. To our knowledge, this is the first report in the cytology literature to describe salivary gland acinic cell carcinoma with metastases to such distant sites and to discuss the resulting cytologic differential diagnoses.

Keywords: Acinic cell carcinoma; Parotid; Cytology; Metastasis; Liver; Vertebral

Abbreviations: AcCC: Acinic Cell Carcinoma; AFP: Alpha-fetoprotein; FNA: Fine Needle Aspiration; HCC: Hepatocellular Carcinoma; PAS-D: Periodic Acid-Schiff Staining with Diastase Digestion; RCC: Renal Cell Carcinoma

Introduction

Salivary gland neoplasms are heterogeneous tumors that vary in anatomic site of origin, histopathology, and behavior, including metastatic potential [1]. While the majority is benign, approximately one-fourth are malignant. Acinic cell carcinoma (AcCC) is a low-grade salivary gland tumor, comprising 1-6% of salivary neoplasms [2-6]. It is slow-growing but capable of metastasizing, with rare reports involving regional lymph nodes, lungs, and bones [7-14]. To date, there have been only few documented cases of vertebral metastases and limited cytologic descriptions of metastatic AcCC. We report a case of recurrent parotid AcCC with widespread liver and bone metastases, diagnosed by fine needle aspiration (FNA) cytology and tissue biopsies.

Case Report

A 40 year old female smoker with a palpable, slowly enlarging left parotid lesion underwent superficial parotidectomy at an outside facility. The tumor was reported to be a completely excised “parotid adenoma” with close adherence to the facial nerve. Three years later, multiple nodules adjacent to the parotidectomy skin flap incision were observed. Repeat excision demonstrated AcCC. In-house review of the initial excision also showed AcCC. The patient subsequently received radiation therapy.

Twelve years after initial parotidectomy, surveillance CT demonstrated a 1.5-cm left parotid bed lesion and lytic lesions in the left fourth rib and T8 vertebra with increased PET uptake. FNA of the lesions showed AcCC. The patient underwent ablation followed by kyphoplasty.

Over the next three years, the patient had widespread metastases to skull, hip, and sacrum, with parotid recurrences. She underwent chemotherapy, completion parotidectomy, and left neck dissection, demonstrating soft tissue invasion by AcCC without regional lymph node involvement.

Eighteen years after initial diagnosis, ultrasound showed echogenic liver lesions measuring up to 1.8 cm. Hypercellular core biopsy touch preparations (Figures 1-4) demonstrated round to polygonal epithelial cells arranged singly and in branching sheets and occasional clusters around capillaries. The cells exhibited relatively monotonous round to oval nuclei with fine chromatin, scattered nucleoli, and abundant granular to focally abundant cytoplasm.

Figure 1: Touch preparation of core needle biopsy of liver lesions (Pap, 40x). Round to polygonal epithelial cells are arranged in branching sheets and clusters around capillaries.

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The background contained bare nuclei, rare lymphocytes, hepatocytes, and granular proteinaceous material. No necrosis, increased mitoses or dedifferentiated areas were observed. The cytomorphologic findings resembled the previous excisions and the corresponding core biopsy confirmed metastatic AcCC (Figure 5). Ablation of liver lesions was performed, and the patient returned to baseline four months later.

**Discussion**

First described by Nasse (1892), AcCC is an uncommon salivary gland neoplasm comprising 7-17.5% of salivary malignancies [15-17]. Eighty percent arise in the parotid gland, with the remainder in the minor salivary, submandibular, and sublingual glands (6%-15%, 4%, and 1%, respectively) [18]. It is the most common bilateral malignant salivary tumor [15].

AcCC presents as a slowly enlarging parotid mass with occasional vague, intermittent pain (40%) and rarely facial muscle weakness or paralysis (<10%). It occurs more frequently in women (female-to-male ratio of 3:2) [5] over a wide age range (mean onset in the fourth decade of life) [6], although cases afflicting children and elderly have been reported [19,20].

Grossly, most AcCCs are solitary, well-circumscribed nodules less than 3 cm in size (range 0.5-13.0 cm) with a firm, rubbery consistency. The cut surface is red-tan and lobular, with occasional solid, cystic or hemorrhagic components [19].

Cytological diagnosis of AcCC is problematic, with a high false-negative rate (~50%) [21]. On FNA, hypercellular smears demonstrate predominantly serous acinar-type cells, which resemble normal salivary gland acini. The polygonal neoplastic cells exhibit bland, round, eccentrically-placed nuclei and a moderate amount of delicate, vacuolated, lightly basophilic cytoplasm with fine to coarse metachromatic zymogen granules. Occasional intranuclear pseudoinclusions can be identified. Several other cell types may be present, including intercalated vacuolated cytoplasm. The background contained bare nuclei, rare lymphocytes, hepatocytes, and granular proteinaceous material.
duct-type cells, nonspecific glandular cells, vacuolated cells, and clear cells. The neoplastic cells may be arranged singly or in sheets and crowded clusters with indistinct cell borders. Other architectural patterns (encountered on histology) include microcystic, papillary-cystic, and follicular [2]. Mitoses, necrosis, and marked cytologic pleomorphism are infrequent. The background contains bare nuclei, vascular-rich stromal fragments, lymphocytes and foamy proteinaceous material. Rarely, ACC shows high-grade transformation/dedifferentiation, in which case the underlying diagnosis is not so obvious. Immunostains are nonspecific. Periodic acid-Schiff staining with diastase digestion (PAS-D) can help highlight the cytoplasmic granules [18].

When ACC metastasizes, the cytologic differential diagnosis must be expanded. In our case with liver metastases, the differential includes a primary hepatic lesion, like hepatocellular carcinoma (HCC). The acinar-type cells of ACC can superficially resemble hepatocytes (Figure 4), due to abundant granular cytoplasm and relatively round nuclei. Intracellular pseudoinclusions may also be seen [15], along with rosette/acinar formations [22].

However, hepatocytes are more "oncocytic," with eosinophilic cytoplasm versus the basophilic, vacuolated to granular cytoplasm seen in ACC. HCC cells occasionally display cytoplasmic bile, Mallory hyaline, or trabecular arrangements surrounded by CD34-positive endothelial cells in a cribriform background [15,23]. Immunostaining for HCC includes hepatocyte antigen (HepPar 1) and arginase-1, as well as polyclonal carcinomaembryonic antigen and CD10 in a canalicular pattern [6,23,24]. PAS-D staining would be more indicative of metastatic ACC.

Other tumors should also be considered in the cytologic differential diagnosis of metastatic salivary ACC to the liver. Primary acinar cell carcinoma of the liver is exceedingly rare, characterized by morphological acinar differentiation, architectural pattern and typical cytological features including pyramidal shape, granular cytoplasm, PAS-D staining, trypsin and chymotrypsin positivity [25,26]. These features resemble salivary ACC, which is reported positive for alpha-1-antichymotrypsin and alpha-1-antitrypsin [14,27]. Primary acinar cell carcinoma of the liver is favored in the absence of another primary site, such as pancreas or salivary glands, and when resection of the liver lesion causes a rapid decrease in serum alpha-fetoprotein (AFP) levels [26]. In our case, the patient had a history of primary parotid ACC and never had serum AFP elevation.

Adrenal cortical carcinoma also displays finely granular chromatin, eccentric nuclei and single cells to loose cell-clusters associated with capillaries. Unlike ACC, it has prominent cytologic atypia, mitoses, necrosis, occasional clear lipid-filled cells, and immunopositivity for synaptophysin, calretinin, inhibin, and Melan A [15].

Occasionally, salivary ACC demonstrates clear cytoplasm, though this rarely predominates on FNA [6]. Regardless of whether the neoplasm is in the salivary gland or distant metastatic sites, the differential diagnosis in such situations includes metastatic clear cell renal cell carcinoma (RCC) [28]. Metastatic RCC may show greater nuclear pleomorphism and glycogen-containing cells, as well as positive immunostaining for CD10, carbonic anhydrase 9, PAX2, PAX8, and RCC [15].

ACC of salivary gland origin can also be mistaken for metastatic papillary thyroid carcinoma [6]. Cellular sheets and follicle-like formations can be seen in both, as well as nuclear pseudo-inclusion [15] and psammoma bodies [6]. Positivity for thyroid transcription factor-1 (TTF-1) and thyroglobulin, and occasional colloid should correctly identify metastatic thyroid carcinoma [15].

ACC is generally considered a low-grade neoplasm with good prognosis (95% survival). However, it has a high recurrence rate (8.3-45%) [8]. Metastatic rate is 2-20% [29]. Metastases usually involve cervical lymph nodes, lungs, and bones, and rarely liver [9,12], skin [14], and orbit [13]. Treatment is complete excision with possible lymph node dissection. Radiation, chemotherapy and ablation may be useful palliatively.

Limited discussion exists in the literature regarding cytologic findings and differential diagnosis of widespread metastases of ACC. There have been only six reports of vertebral ACC metastases [2,8,10,30]. We document this case to not only demonstrate such features and diagnostic considerations when encountering metastatic ACC on FNA, but also to show that despite low-grade histology, there remains the limited possibility of widespread metastasis in patients with a history of ACC.

References


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