Hürthle Cell Carcinoma in an Extraordinary Location: A Case Report

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ABSTRACT

Aim: To present a case of Hürthle cell carcinoma at an extraordinary site, and to contribute to the understanding and management of this tumor.

Background: Hürthle cell carcinomas are a dangerous but uncommon variant of thyroid follicular carcinoma, exhibiting invasive potential, they have a propensity to metastasize to distant sites. Therefore, swift diagnosis and management with multidisciplinary input are warranted. We humbly present a case of Hürthle cell carcinoma in the supraclavicular region.

Case description: A 44-year-old woman, with a prior history of follicular thyroid adenoma, was found to have a mass measuring 20 × 15 × 18 mm on the left supraclavicular region. Magnetic resonance imaging (MRI) and fine needle aspiration biopsy were suspicious of follicular thyroid neoplasm. Positron emission tomography–computed tomography (PET–CT) revealed two lesions with avid fluorodeoxyglucose (FDG)-avid uptake, one corresponding to the original supraclavicular location and one in the left iliac crest. Subsequently, histological analysis following surgical excision of the supraclavicular mass revealed a lobulated Hürthle cell tumor. The patient underwent further cervical lymph node dissection, with one supraclavicular node demonstrating infiltration by Hürthle cell carcinoma. Postoperatively, the patient underwent radioactive iodine (RAI) therapy to which she responded well, and currently remains well at follow-up.

Conclusion: Hürthle cell carcinomas are a rare but aggressive variant of follicular thyroid carcinomas. Known for their malignant potential, this case underscores the vigilance required for the diagnosis and management of these tumors as their sites of metastasis may be most extraordinary, as illustrated in this report. Swift diagnosis and multidisciplinary management of these tumors are paramount.

Clinical significance: Hürthle cell carcinomas have been demonstrated to metastasize to the most extraordinary sites, we present a unique case of Hürthle cell carcinoma which reinforces the notion and hope to contribute to its understanding.

Keywords: Ectopic thyroid tissue, Endocrine surgery, Hürthle cell carcinoma, Oncocytic thyroid carcinoma, Thyroid carcinoma.

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INTRODUCTION

Hürthle cell/oncocytic carcinomas are an uncommon but dangerous variant of follicular carcinoma of the thyroid. They exhibit a wide spectrum of behavior with regards to local invasion, with some variants being highly invasive, infiltrating thyroid parenchyma, and extrathyroidal tissue, while some exhibiting minimally invasive behavior. Furthermore, Hürthle cell carcinomas have a high likelihood of metastasis; hence, their swift diagnosis and treatment are warranted. We herein present an intriguing case of a patient with Hürthle cell variant follicular carcinoma in the ectopic thyroid tissue of the supraclavicular region.

CASE DESCRIPTION

A 44-year-old woman with a history of benign thyroid adenoma presented to her general practitioner (GP) with a 6-month history of a nontender palpable lump in the left supraclavicular region. Her background medical history included a left hemithyroidectomy for a Hürthle cell thyroid adenoma, as well as a hysterectomy for fibroid disease. The lump measured 20 × 15 × 18 mm and was located in the left supraclavicular fossa, 5 cm medial to the acromial-clavicular joint. Ultrasonography of the lesion showed a highly vascular, thin-walled homogeneous mass. A subsequent magnetic resonance imaging (MRI) study confirmed the findings of a highly vascular soft-tissue mass. Fine needle aspiration biopsy of the lesion revealed a diagnosis of Hürthle cell follicular thyroid neoplasm.

A positron emission tomography–computed tomography (PET–CT) showed two fluorodeoxyglucose (FDG)-avid lesions—one in the left supraclavicular fossa and another in the left iliac crest. With radiological findings in hand, the patient underwent an excisional biopsy of the supraclavicular lesion. Histopathology of the excised lesion showed a lobulated Hürthle cell tumor-infiltrating into fibrofatty tissue, with an elevated Ki-67 index, and positive TTF-1 and Pax-8 staining. Biopsies of the adjacent left supraclavicular lymph node also demonstrated lobulated Hürthle cell tumor-infiltrating into fibrofatty tissue, with no lymph nodal tissue or normal thyroidal tissue present.

A right hemithyroidectomy was planned following a multidisciplinary meeting. Histological analysis revealed a normal right hemithyroid. However, tissue from the left thyroid bed showed a focus of Hürthle cell thyroid follicular carcinoma present in adipose tissue and adjacent to skeletal muscle fibers.

Six months after the patient’s initial presentation, a suspicious dermal nodule along with a lymph node on the patient’s left supraclavicular region was identified and excised. The latter was confirmed to be Hürthle cell tumor infiltration upon histological

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analysis. Fortunately, benign nodes were noted in the left level II to V dissection. A consensus arrived at multidisciplinary meetings and radioactive iodine (RAI) therapy was planned for the patient. The patient underwent and responded well to RAI therapy and remained well at clinical follow-up with endocrinology and general surgery.

**Discussion**

Ectopic thyroid tissue in the head and neck, although uncommon, are well-known entities in medical literature, as the development and translocation of the thyroid gland to its physiological paratracheal location have been extensively studied. The reported prevalence of ectopic thyroid tissue is to be at 1 in 100,000 to 300,000 with lingual ectopic sites accounting for 90% of cases. Though elusive, ectopic thyroid tissue in locations, such as, the abdomen have been reported. We believe that this is the first reported case of metastatic Hürthle cell thyroid carcinoma of the supraclavicular fossa in the medical literature.

Hürthle cell carcinomas encompass approximately 5% of all differentiated thyroid carcinomas. The paucity of its prevalence, aggressive clinical behavior, along with the difficulty in distinguishing metastatic disease of this variant contributes to the ongoing debate regarding its clinical course and prognosis. Hürthle cell carcinomas are histologically subdivided into widely invasive and minimally invasive neoplasms. Despite its classical appearance, it is impossible to definitively diagnose malignancy based solely on cytological features; Hürthle cell lesions traditionally require surgical excision and biopsy for definitive diagnosis. The fact that even with extensive pathological evaluation, some Hürthle cell neoplasms will remain classified as indeterminate behavior reinforces the intricacy of these tumors.

While there is medical literature on the treatment of Hürthle cell carcinomas of the thyroid, the literature on the treatment of follicular carcinoma in ectopic thyroid tissue is alarmingly scarce. It would not be a stretch to consider the management of our patient’s case to be uncharted territory in medicine; the current paucity in its treatment of Hürthle cell carcinoma in ectopic thyroid tissue is alarmingly scarce. It would not be a stretch to consider the management of our patient’s case to be uncharted territory in medicine; the current paucity in its treatment of Hürthle cell carcinoma in ectopic thyroid tissue is alarmingly scarce.

**Conclusion**

We sought to present this case of a rare Hürthle cell follicular thyroid carcinoma in an extraordinary anatomical site of the left supraclavicular fossa, which we believe is the first to be reported. We hope that we were able to contribute to the current and future understanding of this rare and often aggressive tumor.

**Clinical Significance**

Hürthle cell carcinoma is a very uncommon tumor, but with its aggressive nature, such malignancies must be promptly diagnosed and treated. In this report, we have described a case of Hürthle cell carcinoma in an extraordinary location, in doing so we hope to contribute to the current body of knowledge on the management of such a tumor.

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