Unusual association between adrenal leiomyoma and autoimmune disease

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Adrenal leiomyoma is a rare benign deep soft tissue tumor, which arises from the residual embryonal blood vessel tissue, and smooth muscle fibers. Few cases were reported in the literature, most of which were non-functioning, incidentally discovered, and associated with immune deficiency. On literature review, none of the reported cases were associated with a concurrent autoimmune disease. Herein, we report a rare case of adrenal leiomyoma in association with Hashimoto's thyroiditis and review the literature.

A 38-year-old Mediterranean woman, a known case of Hashimoto’s thyroiditis, presented to Jordan University Hospital, Amman, Jordan, with signs of complicated acute appendicitis. A CT-scan was performed and showed an incidental right adrenal mass measuring 3 x 2 x 2 cm. The adrenal mass was investigated after appendectomy and proved to be non-functioning. The other adrenal gland was normal. She had normal blood pressure. Urine 3-methoxy-4-hydroxymandelic acid (VMA) and metanephrine levels were within normal limits. Serum electrolytes and cortisol levels (dexamethasone suppression test) were within normal limits. Human immunodeficiency virus immunoglobulin-G (HIV IgG) titer was negative.
Leiomyomas are benign tumors consisting of smooth muscle cells. They occur most frequently in the uterus and gastrointestinal tract, but have been found in a wide range of sites including the adrenal glands. A malignant counterpart is seen in the adrenal gland, but is exceedingly rare. Adrenal leiomyomas are usually small tumors that are not highly worrisome for adrenocortical carcinoma. While the National Institutes of Health (NIH) recommends surgical resection of incidental adrenal tumors larger than 6 cm in greatest dimension, most of the reported cases of adrenal leiomyomas were smaller than that size. Fluorodeoxyglucose-positron emission tomography (FDG-PET) scan revealed that the tumor was non-functioning in most reported cases, with unequivocal results in the remaining ones.

Interestingly, patients with acquired immunodeficiency syndrome (AIDS) especially children, are at an increased risk for the development of smooth muscle neoplasms, such as leiomyoma and leiomyosarcoma. There is an association between

Thyroid function tests were normal with L-thyroxin treatment. Anti-thyroid peroxidase antibodies (TPO) were elevated (239 IU; <150 IU/ml [normal range]), anti-thyroglobulin titer was negative (less than 85 IU, <85 IU [normal range]). The initial diagnosis of Hashimoto’s thyroiditis was based on clinical and serologic tests. A repeat CT scan 6 months later showed enlargement of the mass, with a new measurement of 4 x 4 x 3 cm. She underwent successful right adrenalectomy (Fig re 1). Histologically, the adrenal gland was totally occupied by a tumor, composed of uniform spindle cells arranged in interlacing fascicles. No nuclear pleomorphism, coagulative necrosis, or increased mitotic activity was seen (Fig re 2). Dense lymphocytic infiltrate was noted in the background of the tumor in all levels, both intratumoral and peritumoral (Fig re 3). Immunohistochemical studies confirmed the diagnosis of adrenal leiomyoma (Fig res 4 & 5). There were no post-operative complications, and the patient was discharged home 4 days after surgery.
Adrenal leiomyoma and latent Epstein Barr virus infection (EBV), and/or HIV/AIDS. Five among the reported cases were associated with HIV/AIDS, and one with ataxia-telangiectasia. An EBV infection is associated with smooth muscle tumors in immunocompromised, but not in immunocompetent patients. The patient in this paper was seronegative for HIV. The EBV IgG antibody was positive, but IgM was negative. Immunohistochemical stain for EBV in the histological specimen was negative. On literature review, none of the reported cases showed an association with an autoimmune process, or an additional peculiar histopathological feature, as in our case. Based on the histopathological finding in our case, concurrent Hashimoto’s thyroiditis, and being a middle aged female patient, we suppose an association between the adrenal leiomyoma and an autoimmune process. This observation can be noteworthy, especially if it is reported again in the future.

We conclude that adrenal leiomyoma is rare and is associated with good prognosis. The local recurrence and transformation into sarcoma is rare. Surgeons, pathologists, and endocrinologists should be aware of the possibility of an adrenal leiomyoma as a cause of adrenal incidentalomas, and complete excision is recommended in these cases.


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