Retrophreitoneal liposarcoma mimicking pheochromocytoma

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ABSTRACT

Retrophreitoneal liposarcomas are rare mesenchymal tumors which carry a poor prognosis because of their late presentation. When symptoms do occur, they are usually nonspecific and related to mass effect or invasion of local structures. Rarely, retrophreitoneal liposarcomas can clinically and biochemically mimic pheochromocytomas. We discuss one such case of a 56-year-old Afro-Trinidadian female who presented to her primary care physician with a 3-month history of weakness, intermittent sweating, difficulty sleeping and elevated blood pressure. After a 2 week trial of an oral antihypertensive regime her blood pressure was still elevated and she complained of new right sided abdominal pain. A subsequent Computed Tomography scan of her abdomen revealed an enhancing, heterogeneous right suprarenal mass suspicious for pheochromocytoma. Urinary catecholamines were also elevated and an MRI of her abdomen supported the diagnosis of pheochromocytoma although intrasional fat was noted, an uncommon feature of pheochromocytomas. She was booked for laparoscopic adrenalectomy. Histological analysis of the resected specimen confirmed a dedifferentiated retrophreitoneal liposarcoma. While the imaging features of pheochromocytomas and retrophreitoneal liposarcomas can be similar, the presence of intrasional fat on CT and MRI should favour the diagnosis of a retrophreitoneal liposarcoma, albeit the clinical and biochemical picture.

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**Introduction**

Liposarcomas are malignant tumors of mesenchymal origin that can arise from any fat containing region, including the retroperitoneum. They are the most common sarcoma of the retroperitoneum with a frequency of 41% followed by leiomyosarcoma and malignant fibrous histiocytoma [1]. In the retroperitoneum about 35% arise in the perinephric fat [2]. Histologically, liposarcomas are divided into five types including: well-differentiated, dedifferentiated, myxoid/round cell, pleomorphic and mixed. The well-differentiated type occurs most frequently while the dedifferentiated and pleomorphic types are aggressive with metastases often seen at presentation. Rarely, they may mimic a phaeochromocytoma. We contribute one more such case to the literature.

**Case report**

A previously well 56-year-old Trinidadian female of African descent presented to her primary care physician with a three-month history of weakness, intermittent sweating and difficulty sleeping. Her blood pressure was found to be elevated (usually above 170/100 mm Hg) and she was started on enalapril 10 mg and nifedipine sustained release 20 mg daily. Two weeks later, her blood pressure were still uncontrolled and remained markedly elevated, and there was a new complaint of right upper quadrant abdominal pain. Physical exam was unremarkable with no palpable abdominal masses. A Computed Tomography (CT) scan of her abdomen and pelvis with IV contrast was requested which demonstrated a heterogeneous, avidly enhancing, right suprarenal mass (Figs 1 and 2). An MRI of her abdomen and urine catecholamines (24-hour 5-hydroxyindoleacetic acid [5-HIAA] and 24-hour metanephrine) were requested to investigate for possible pheochromocytoma. The MRI revealed an intermediate T1, mildly hyperintense T2, right suprarenal mass with heterogeneous enhancement post IV gadolinium (Figs. 3, 4 and 5). Foci of signal dropout were visualized on the out of phase sequences consistent with intrallesional fat (Fig. 6). The right adrenal gland was seen in close relation to this mass. Urinary metanephrine was marginally elevated at 218 microgram/24 h (reference range: 36-209 microgram/24 h) as was 5-HIAA at 22.7 mg/24 h (reference range: 0-14.9 mg/24 h). Though the presence of intrallesional fat is uncommon for pheochromocytomas, the clinical presentation and elevated urine catecholamines supported pheochromocytoma as the primary differential diagnosis. The patient was referred for preoperative medical management with an alpha blocker (terazosin 2 mg nocte), followed by beta blocker (propranolol 10 mg bid) with good effect such that her blood pressure rarely exceeded 125/75 mmHg in the 2-week interval from surgery. Laparoscopic adrenalectomy was then performed with initial control of the right adrenal vein. There was no typical drop in the blood pressure once this was done. Slight bleeding was encountered laterally and superiorly with subsequent removal of the perinephric fat pad for control. The patient was discharged on post-op day 1. Histological examination of the resected specimen revealed a dedifferentiated liposarcoma (Fig. 7) with positive resection margins. There was no histological evidence of pheochromocytoma and the right adrenal gland was normal. Interestingly, there was no further need for antihypertensive medications even after 6 weeks of having surgery and her abdominal pain.

**Fig. 1 – Axial non contrast CT image showing a well circumscribed soft tissue density right suprarenal mass (white arrow) containing a few tiny hypodense foci within (yellow arrows). No calcifications were visualised (colour version of figure is available online).**
Fig. 2 – Arterial phase post IV contrast CT imaged demonstrating marked hypervascularity of the right suprarenal mass (white arrow), which is separate from the posterior segment of the right lobe of liver and the right adrenal gland (black arrow).

Fig. 3 – Axial T2 weighted MRI image showing a well-circumscribed, heterogenous, mildly hyperintense right suprarenal lesion (white arrow).

resolved. A post-op CT demonstrated a non-enhancing right suprarenal soft-tissue density, suggestive of post-operative changes (Fig. 8).

Discussion

Retroperitoneal liposarcomas are a subtype of liposarcomas originating from retroperitoneal fat. They occur at any age but tend to occur most frequently in the sixth and seventh decades of life with no gender or sex predilection [3]. The large potential space of the retroperitoneum allows retroperitoneal liposarcomas to grow quite big before presenting with symptoms. When symptoms do occur, they are usually non-specific and include abdominal and flank pain, early satiety, and lower limb swelling and pain [4]. Neurological, muscularkeletal and obstructive symptoms can occur secondary to local invasion or compression of adjacent structures [4]. Our presentation of a retroperitoneal liposarcoma mimicking a pheochromocytoma with markedly elevated blood pressures, elevated urine catecholamines and suggestive imaging features is uncommon but previously documented by Marmouch H et al. [5]. Dedifferentiated liposarcomas have been shown to present with elevated catecholamines, and demonstrate somatostatin receptors both on imaging (somatostatin receptor
Fig. 4 – Coronal T2 weighted MRI image demonstrating the relationship of the right suprarenal mass (white arrow) to the right kidney (red arrow), right adrenal gland (black arrow) and the right lobe of the liver (colour version of figure is available online).

Fig. 5 – Axial T1 weighted post IV gadolinium MRI image demonstrating hypervascularity and heterogeneous enhancement within the right suprarenal mass (white arrow).

scintigraphy) and histopathology (reverse transcription polymerase chain reaction) [6].

CT is used for the diagnosis, staging and preoperative evaluation of liposarcomas. On CT, these tumors contain fat and varying degrees of soft tissue. Low grade liposarcomas contain mostly fat and very little soft tissue, intermediate-grade tumors are relatively lucent with transverse septations and higher-grade sarcomas are dense, heterogenous and enhance with IV contrast [4]. Our case of a dedifferentiated liposarcoma appeared as a heterogenous, enhancing soft tissue density mass with a few small fatty foci. Calcifications were not present but when visualized are associated with a poor prognosis [4]. CT can also be used to determine invasion of adjacent structures, assess regional lymph node status and determine the presence of liver and lung metastases. MRI is used in cases of an indeterminate retroperitoneal mass and to also determine neurovascular invasion. Abdominal ultrasound when done, demonstrates a hyperechoic mass but fails to separate retroperitoneal liposarcomas from other retroperitoneal masses. CT and ultrasound may be used to guide percutaneous biopsy but histopathological examination of the surgical specimen remains the gold standard [7,8].

Management of retroperitoneal liposarcomas is wide local excision with negative resection margins. En bloc resection can however be challenging when the tumor encases vital structures such as the great vessels. A balance between removing tumor and patient safety should therefore form the basis of surgical management in such cases. Attempts at
neoadjuvant chemotherapy have proven useless [9]. Haas RL et al. concluded that perioperative radiotherapy has been associated with better local control but level I evidence is still required [10]. An open approach is generally preferred by most surgeons because of tumor bulk and the need to removal adjacent organs. Today with growing expertise in minimally invasive surgery, a laparoscopic approach can be used in very selected patients with limited disease which would offer reasonable vision of the surgical field, less post-operative pain and better cosmesis [11]. Poor prognosis is seen in cases with a positive resection margin, higher-grade tumors, and multifocal disease. High-grade tumors are also associated with higher local recurrence rates and distant metastases. The recurrence rate of dedifferentiated tumors is about 80% [9,12]. The five-year survival rate of low-grade well differentiated and myxoid tumors is 90% while the 5 year survival rate of high-grade pleomorphic, round cell and dedifferentiated tumors is 30%-50%, 60% and 75% respectively [4]. Our patient will receive a follow up MRI scan in 3 months and further surgery determined as necessary.

In conclusion, although unusual, retroperitoneal liposarcomas can rarely mimic a phaeochromocytoma. Typically retroperitoneal liposarcomas produce symptoms, when large, secondary to mass effect and invasion of adjacent struc-
ures. While imaging features of phaeochromocytomas and retroperitoneal liposarcomas can be similar, the presence of intrallesional fat on CT and MRI should favour diagnosis of the latter, despite the clinical and biochemical picture.

**Patient consent**

Informed consent was obtained from the patient to use her information in this manuscript.

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