Case Report

Adrenal Lymphangioma Masquerading as a Catecholamine Producing Tumor

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Objective. To report the unusual case of an adrenal lymphangioma presenting in a patient with an adrenal cystic lesion and biochemical testing concerning for pheochromocytoma. The pertinent diagnostic and imaging features of adrenal lymphangiomas are reviewed.

Methods. We describe a 59-year-old patient who presented with hyperhidrosis and a 2.2 by 2.2 cm left adrenal nodule. Biochemical evaluation revealed elevated plasma-free normetanephrine, urine normetanephrine, urine vanillylmandelic acid, and urine norepinephrine levels. Elevated plasma norepinephrine levels were not suppressed appropriately with clonidine administration.

Results. Given persistent concern for pheochromocytoma, the patient underwent adrenalectomy. The final pathology was consistent with adrenal lymphangioma.

Conclusions. Lymphangiomas are benign vascular lesions that can very rarely occur in the adrenal gland. Imaging findings are generally consistent with a cyst but are nonspecific. Excluding malignancy in patients presenting with adrenal cysts can be difficult. Despite its benign nature, the diagnosis of adrenal lymphangioma may ultimately require pathology.

1. Introduction

Lymphangiomas are benign vascular lesions that most commonly occur in the head, neck, and axilla [1]. Lymphangiomas of the adrenal gland are very rare, with an estimated incidence of 0.064 to 0.18 percent in autopsy series [2]. These lesions are generally asymptomatic but may be found incidentally on radiographic imaging during work-up for unrelated conditions. Therefore, it is important to distinguish these benign cysts from malignant adrenal lesions. Here, we present the diagnostic dilemma of a patient who presented with an adrenal cystic lesion and biochemical evaluation concerning for pheochromocytoma. The patient was ultimately found to have an adrenal lymphangioma.

2. Case Report

A 59-year-old male with past medical history of hypertension, obesity, hyperhidrosis, and secondary polycythemia presented to his outpatient hematologist in 2009 for evaluation of hyperhidrosis. The patient endorsed several-year history of profuse episodic sweating that interfered with his social interactions. He denied any headaches or palpitations. The patient had a known history of 1.7 by 1.4 cm adrenal nodule discovered incidentally on MR scan performed that same year. The nodule did not enhance, and imaging characteristics were most consistent with a cyst.

Biochemical evaluation in 2009 included an elevated 24-hour urine normetanephrine, norepinephrine, and vanillylmandelic acid as indicated in Table 1. Urine 24-hour metanephrine and epinephrine were both within normal limits. The patient was not taking any medications that could interfere with the measurement of catecholamines or metanephrines. In 2010, the patient underwent an 11131I metaiodobenzylguanidine (MIBG) scan that was negative for pheochromocytoma.

Given continued symptoms of hyperhidrosis, the patient was referred to endocrinology in 2013. Repeat biochemical
Table 1: Patient’s catecholamine levels before and after left adrenalectomy.

<table>
<thead>
<tr>
<th></th>
<th>2009 (Preoperative)</th>
<th>Reference range</th>
<th>2013 (Postoperative)</th>
<th>Reference range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plasma-free normetanephrine</td>
<td>nd</td>
<td>nd</td>
<td>1.6 nmol/L</td>
<td>&lt;0.90 nmol/L</td>
</tr>
<tr>
<td>Plasma-free metanephrine</td>
<td>nd</td>
<td>nd</td>
<td>0.53 nmol/L</td>
<td>&lt;0.90 nmol/L</td>
</tr>
<tr>
<td>Plasma norepinephrine</td>
<td>nd</td>
<td>nd</td>
<td>0.50 nmol/L</td>
<td>nd</td>
</tr>
<tr>
<td>Plasma epinephrine</td>
<td>nd</td>
<td>nd</td>
<td>0.20 nmol/L</td>
<td>nd</td>
</tr>
<tr>
<td>Urine normetanephrine</td>
<td>1868 µg/24 hours</td>
<td>110–1050 µg/24 hours</td>
<td>1123 µg/24 hours</td>
<td>50–800 µg/24 hours</td>
</tr>
<tr>
<td>Urine metanephrines</td>
<td>254 µg/24 hours</td>
<td>35–460 µg/24 hours</td>
<td>123 µg/24 hours</td>
<td>0–300 µg/24 hours</td>
</tr>
<tr>
<td>Urine norepinephrine</td>
<td>414 µg/24 hours</td>
<td>0–140 µg/24 hours</td>
<td>257 µg/24 hours</td>
<td>0–100 µg/24 hours</td>
</tr>
<tr>
<td>Urine epinephrine</td>
<td>16 µg/24 hours</td>
<td>0–32 µg/24 hours</td>
<td>10.2 µg/24 hours</td>
<td>0–20 µg/24 hours</td>
</tr>
<tr>
<td>Urine vanillylmandelic acid</td>
<td>9.6 µg/24 hours</td>
<td>0.0–7.5 µg/24 hours</td>
<td>nd</td>
<td>nd</td>
</tr>
</tbody>
</table>

nd: not done.
Results outside the reference range are indicated in bold.

evaluation at that time was notable for an elevated plasma-free normetanephrine, plasma norepinephrine, 24-hour urine normetanephrine, and 24-hour urine norepinephrine as documented in Table 1. Plasma-free metanephrine, plasma epinephrine, urine 24-hour metanephrine, and urine 24-hour epinephrine were within normal limits. He was also diagnosed with new onset diabetes.

With regard to further imaging studies, adrenal protocol CT demonstrated a 2.1 by 2.2 cm left adrenal nodule which could not be classified as lipid rich adenoma. The density measurement of the nodule was 12 Hounsfield units before contrast and 22 Hounsfield units on the enhanced study. The absolute enhancement washout value was 20%, below the 60% threshold for an adenoma. Adrenal protocol MR demonstrated a 2.2 × 2.2 cm lesion with uniformly high signal intensity on T2-weighted sequences and low signal on T1-weighted images with no enhancement on contrast administration. MR findings were most consistent with a cyst (Figure 1).

Given the patient’s modestly elevated normetanephrine and norepinephrine, confirmatory testing was performed with a clonidine suppression test. Norepinephrine was 891 pg/mL (normal reference range <500 pg/mL) prior to 0.3 mg clonidine administration and 708 pg/mL 3 hours after clonidine administration. Normetanephrine was 0.95 nmol/L (normal reference range <0.90 nmol/L) prior to clonidine administration and 0.94 nmol/L 3 hours after clonidine administration which is suggestive of pheochromocytoma. Due to persistent concern for potential pheochromocytoma with cystic degeneration, he was started on phenoxycbenzamine and underwent laparoscopic left adrenalectomy without complications. Surgical pathology demonstrated a benign vascular cyst consistent with lymphangioma (Figure 2).

Repeat plasma fractionated metanephrines performed after surgery were within normal limits. Incidentally, patient reported significant improvement in his hyperhidrosis after surgery and also significant weight loss and improvement of diabetes control. Polycthyemia was most likely not related to the adrenal tumor, since no changes have been noticed up to 6-month postop follow-up.

3. Discussion

Adrenal cysts can be classified as pseudocysts, endothelial cysts, epithelial cysts, or parasitic cysts. The estimated frequency of each subtype varies by series, with approximately 39–78% classified as pseudocysts, 20–45% endothelial cysts, 2–9% epithelial cysts, and 0–7% parasitic cysts [3–5]. Endothelial cysts can be further divided into lymphangiomaticous and angiomatous cysts. Lymphangiomas are believed to arise from faulty lymphatic development leading to either isolation of the lymphangioma from larger lymphatic channels or lack of fusion with the venous system [1, 4, 6].

An estimated 7–15% of adrenal cysts are associated with malignancy [3, 5, 7, 8]. Although lymphangiomas are benign lesions, aspiration or surgical removal becomes necessary when malignancy cannot be excluded based on imaging alone. On ultrasonography, lymphangiomas generally present as well-demarcated multiloculated cysts. Less commonly, they present as unicameral cysts [9, 10]. On CT scan, they display capsular enhancement with internal attenuation values generally in the range of water [9–11]. On MR, these cysts are hypointense and nonenhancing on T1-weighted images with no enhancement on contrast administration. MR findings were most consistent with a cyst (Figure 1).

Identification of surgically excised adrenal cysts can be confirmed histologically. Adrenal lymphangiomas are characterized by multicystic architecture with a simple endothelial lining. On immunohistochemistry, these lesions stain negative for keratin and positive for D2-40, a marker of lymphatic endothelium [4].
tumor. Given the low prevalence of pheochromocytoma in the general population, even screening tests with high specificity will have more false-positive results than true-positive results [12, 13]. In cases where there is high clinical suspicion for pheochromocytoma, plasma-free metanephrines and normetanephrines are an appropriate first screening test given their high reported sensitivity of 96–99% [13,14]. Large elevations in plasma metanephrines or normetanephrines several times the upper limit of normal are highly suggestive of pheochromocytoma and should prompt further investigation to localize the tumor [12–15]. In cases where plasma metanephrine or normetanephrine elevations are mild, further testing can be performed to help confirm or exclude the diagnosis including plasma catecholamines, urinary fractionated metanephrines, urinary catecholamines, and vanillylmandelic acid. Factors that can interfere with the diagnostic accuracy of these tests should also be addressed. Physiologic stress, obstructive sleep apnea, caffeine, nicotine, and several medications including many antihypertensives, antidepressants, stimulants, and sympathomimetics can all interfere with test results and lead to false-positive results [13].

In cases where biochemical test results are equivocal, a clonidine suppression test can be used to help clarify the diagnosis. Clonidine is a central alpha2-agonist that suppresses catecholamine release by the sympathetic nervous system. Release of catecholamines from a pheochromocytoma is thought to be autonomous and therefore would not be suppressed by clonidine [16]. Although several criteria have been proposed to define an appropriate clonidine response, a 3-hour postclonidine plasma norepinephrine level of less than 500 pg/mL with a decline of at least 50% has been used with a sensitivity and specificity of 97% and 74%, respectively [17]. The specificity of the test is significantly reduced in patients with a preclonidine plasma norepinephrine level in the normal range [18]. Yet, even using the much more specific criteria which define an abnormal test result simply as a plasma norepinephrine level of 500 pg/mL or less after clonidine administration (specificity of 96%) [17], our patient
still had an abnormal clonidine response. Beta-blockers, tricyclic antidepressants, and thiazide diuretics have all been reported to produce false-positive results; however, none of these medications are applicable to our patient. This case illustrates the difficulty in ruling out pheochromocytoma in patients with benign adrenal lesions.

Adrenal cysts are usually asymptomatic, although local symptoms can vary with the size and position of the lesion. In one case series of 9 patients with adrenal lymphangioma 44% (4/9) presented with abdominal, flank, or back pain while an additional 44% (4/9) were asymptomatic and found incidentally [4]. In the same series one case was found on work-up for labile hypertension that reportedly normalized after resection [19]. Interestingly, our patient reported resolution of his hyperhidrosis after surgery. However, given that adrenal lymphangiomas are nonfunctional cysts, the mechanism by which this could be tied to his hyperhidrosis remains unclear.

4. Conclusion

Here, we present the diagnostic dilemma of a patient with a cystic adrenal lesion in the setting of laboratory testing concerning for pheochromocytoma. Work-up included an elevated plasma norepinephrine level that was not suppressed appropriately with clonidine administration. The patient ultimately underwent adrenalectomy that revealed an adrenal lymphangioma. This case illustrates the difficulty of definitely excluding pheochromocytoma in a patient with a rare benign adrenal cyst.

Conflict of Interests

Dr. Andreas G. Moraitis was previously faculty at the University of Michigan, and currently he works as a medical director at Corcept Therapeutics.

References
