An Unusual Case of Aldosterone- and Norepinephrine-Secreting Retroperitoneal Leiomyosarcoma

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Case Report

A 38-year-old woman presented symptoms of headache, nocturia and fatigue for 4 months. After admission to the hospital, her blood pressure was 190/130 mm Hg, and her pulse was 120 beats/min. A painless, non-pulsatile, firm abdominal mass was palpable at the middle right abdomen, but no other positive sign was found. Serum biochemistry and hormone tests showed: potassium 2.72 mmol/l, norepinephrine 708 ng/ml, adrenaline <50 ng/ml, and aldosterone in the supine position 452.09 ng/l. Computed tomography (CT) and ultrasound of the abdomen showed normal kidneys and adrenal glands, but a large mass was detected in the middle right retroperitoneal cavity (fig. 1). The mass was measured 5.3 cm in vertical extension, 6.4 cm in transverse diameter, and 4.8 cm in anteroposterior diameter. The patient was diagnosed as being a LMS by pathologic examination. We suggested that this was the first case of retroperitoneal LMS shown to be associated with increased levels of serum aldosterone and norepinephrine.

Key Words
Ectopic hormone production · Leiomyosarcoma · Aldosterone · Norepinephrine

Abstract
A case of retroperitoneal leiomyosarcoma (LMS) in a 38-year-old woman who had a tumor in the retroperitoneum with severe hypertension and hypokalemia is presented. Further investigation revealed an elevated serum aldosterone and norepinephrine level. After tumorectomy, the levels of serum aldosterone and norepinephrine successfully normalized; hypertension and hypokalemia were also cured. The tumor was diagnosed as being a LMS by pathologic examination. We suggested that this was the first case of retroperitoneal LMS shown to be associated with increased levels of serum aldosterone and norepinephrine.

Introduction
Leiomyosarcoma (LMS) of the retroperitoneum, belonging to the soft-tissue sarcomas, is relatively uncommon and comprises less than 1% of all malignancies [1]. Usually it has non-specific clinical signs and symptoms. The ectopic hormones secreting LMS is uncommon, especially ectopic catecholamine or aldosterone producing LMS is extremely scarce, and there are no reports. We describe a patient with unusual aldosterone- and norepinephrine-producing retroperitoneal LMS, which was diagnosed by histological and immunohistochemical examination after tumorectomy.

Case Report
A 38-year-old woman presented symptoms of headache, nocturia and fatigue for 4 months. After admission to the hospital, her blood pressure was 190/130 mm Hg, and her pulse was 120 beats/min. A painless, non-pulsatile, firm abdominal mass was palpable at the middle right abdomen, but no other positive sign was found. Serum biochemistry and hormone tests showed: potassium 2.72 mmol/l, norepinephrine 708 ng/ml, adrenaline <50 ng/ml, and aldosterone in the supine position 452.09 ng/l. Computed tomography (CT) and ultrasound of the abdomen showed normal kidneys and adrenal glands, but a large mass was detected in the middle right retroperitoneal cavity (fig. 1). The mass was measured 5.3 cm in vertical extension, 6.4 cm in transverse diameter, and 4.8 cm in anteroposterior diameter. The patient was diagnosed as ‘ectopic pheochromocytoma’, and treated with phenoxybenzamine and potassium chloride. Two weeks later, the hypertension was controlled and serum potassium was normal, and the tumor was subsequently resected. After tumorectomy, the tests of serum biochemistry recovered to normal limits: potassium...
um 3.99 mmol/l, norepinephrine 119 ng/ml, and aldosterone in the supine position 122.7 ng/l. Moreover, hypertension and hypokalemia were cured without application of hypotensive drugs and potassium chloride.

Macroscopically, the tumor mass measured 7 × 6 × 4.5 cm, was well-circumscribed, lobular, and grey-white in color. On the cut surface the tumor was white as well as firm with foci of yellow and soft areas. Microscopically, the tumor was composed of spindle-shaped cells with abundant eosinophilic cytoplasm and elongated nuclei (fig. 2a, b). Most nuclei were centrally located and blunt-ended and had a 'cigar-shaped' appearance (fig. 2b). In some areas of the tumor, nuclear hyperchromatism and pleomorphism were notable and multinucleated giant cells were also readily seen. The spindle-shaped cells, slender or slightly plump, were arranged in fascicles intersecting at right angles (fig. 2a). The tumor was compactly cellular, but myxoid change was focally present with a reticular hypocellular appearance. The coagulative necrosis was evident. Mitotic figures were over 15/10 high-power fields (HPF), and atypical mitoses were often seen. Immunohistochemically, tumor cells were diffusely and strongly positive for desmin, SMA and caldesmon, focally positive for CD34 and EMA, while S-100 and CD117 were negative (fig. 2c–f). The percentage of positive tumor cells for Ki-67 was about 25%. Thus, the tumor was diagnosed as LMS.

Discussion

LMS of soft tissue, usually located in the retroperitoneum, is a relatively unusual mesenchymal tumor that exhibits smooth-muscle differentiation. It showed both a typical cytologic feature, such as spindle-shaped cells with eosinophilic cytoplasm and elongated 'cigar-shaped' nuclei, and an architectural feature, a fascicular growth pattern. Nevertheless, these features of soft-tissue LMSs...
may also occur in other spindle cell tumors, such as myofibroblastic tumors, fibrosarcomas, solitary fibrous tumors (SFTs), malignant peripheral nerve sheath tumors (MPNSTs), extragastrointestinal stromal tumors (EGISTs), dedifferentiated liposarcomas, leiomyomas, and so on. Mostly, the cytologic and architectural features mentioned above could play an important role in the differential diagnosis. Besides these features, in most instances, immunohistochemistry may provide another useful clue to the differential diagnosis. The immunoreactivity of desmin, a specific marker of muscle cell differentiation, is rarely detectable in myofibroblastic tumors and fibrosarcomas, but is encountered, usually strongly and diffusely, in myogenic neoplasms, such as...
LMSs. MPNSTs do not show immunoreactivity for desmin, but may display variable S-100 positivity. SFTs characteristically express CD34 and CD99, but do not or only focally and limitedly express desmin [2, 3]. Most EGISTs are positive for CD117 (c-Kit, a receptor tyrosine kinase encoded from the c-kit proto-oncogene) and CD34, and only less than 5% of EGISTs show reactivity for desmin focally [4]. Dedifferentiated liposarcoma is composed of spindle cells and may show myogenic differentiation as being desmin-positive, but it is usually a typically abrupt transition between the well-differentiated liposarcoma and non-lipogenic area. Thus, both the histologic and immuophenotypic features reliably allow LMSs to differentiate from other spindle-shaped tumor cells. Differentiation between leiomyomas and LMSs of soft tissue may be difficult. Histological assessment of malignancy includes mitotic activity, cellularity, atypia and necrosis. The mitotic activity is the main criterion, namely those tumors with >5 mitoses per 10 HPF are considered malignant. Nuclear antigen Ki-67, which is associated with cell proliferation, is found to be significantly elevated in LMSs, and could be used as an immunohistochemical marker to distinguish between LMSs and leiomyomas [5]. In this case, the tumor shows as histologic features spindle-shaped cells with abundant eosinophilic cytoplasm, elongated nuclei and >15 mitoses per 10 HPF (fig. 2a). The immunoreactivity characteristics of the tumor were strongly and diffusely expression of desmin, SMA and caldesmon, but negative of S-100 and CD117 (fig. 2c–f). Therefore, the tumor was diagnosed as LMS.

Typically, retroperitoneal LMSs have vague presenting symptoms. Moreover, LMSs have rarely produced ectopic hormones, such as β-HCG [6–8], insulin-like growth factor 2 [9], parathyroid hormone [10], and renin [11–13]. Hormone-secreting LMSs may originate from the small intestine, transverse colon, spermatic cord, thorax, or chest wall [7]. There were some case reports of retroperitoneal LMS which secreted renin and caused severe hypertension [11–13]. In view of the high norepinephrine and aldosterone serum level in this patient and the normal limit of hormones after tumorectomy, we believed that this tumor was associated with producing norepinephrine and aldosterone which led to the clinical manifestation of hypertension and hypokalemia. In order to declare the characteristics of the tumor, histological and immunohistochemical examinations were performed, and LMS was diagnosed. To our knowledge, this is the first case report to describe aldosterone- and norepinephrine-secreting retroperitoneal LMS – a new kind of ectopic hormone producing LMS.

Frequently, aldosterone- and norepinephrine-secreting tumors originate from the adrenal gland. Ectopic norepinephrine-secreting tumors are unusual, except ectopic pheochromocytomas. Uede et al. [14] described a case of malignant mixed mesodermal tumor of the uterine corpus with hypercatecholaminemia. It seemed to show that a tumor of mesenchymal origin may produce catecholamine, such as norepinephrine and adrenaline. Ectopic aldosterone-producing tumors are scarce; only a few cases, located in the retrocaval region [15] and ovaries [16, 17], have been described. Up to now, no aldosterone-producing LMS has been presented. This is the first case of retroperitoneal LMS shown to be associated with increased levels of serum aldosterone and norepinephrine. Moreover, it has reminded us that LMS, as a kind of tumor of mesenchymal origin, has an extensively variable ectopic hormone production and clinical manifestation.

References


