CASE REPORT

Primary leiomyosarcoma of the adrenal gland

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Abstract

We report a rare case of a primary leiomyosarcoma of the adrenal gland. A 73-year-old woman presented with an inferior vena cava syndrome. MR imaging was suggestive of a large tumour originating from the right adrenal gland. Angiography revealed a tumour vascularised by the right adrenal artery. At explorative laparotomy a tumour of 27 cm in diameter was found which was completely fixed to the liver; the tumour was therefore considered unresectable. As a consequence of the mechanical problems caused by this large tumour, the patient died 3 weeks after the operation. Autopsy revealed no distant metastases or other primary tumour site.

Key words: leiomyosarcoma, adrenal, angiography

Introduction

Primary mesenchymal neoplasms of the adrenal gland constitute a heterogeneous group of rare entities, including myelolipoma, haemangioma, lymphangioma, angiosarcoma, neurilemoma, leiomymoma and leiomyosarcoma.\textsuperscript{1} A primary adrenal leiomyosarcoma is exceptionally rare, with only four documented cases available in the literature.\textsuperscript{2–5} Here we describe a case, that presented with an inferior vena cava syndrome. Based on detailed angiography of the arterial supply of the tumour, as well as the autopsy data, the tumour was classified as originating from the right adrenal gland. The histological features showed characteristics of leiomyosarcoma.

Case report

A 73-year-old woman was referred to our hospital for further evaluation and assessment of surgical therapeutic options of an intra- or retroperitoneal soft tissue tumour, which caused inferior vena cava syndrome. Three months prior to admission, the patient had noticed swelling of both legs, and she experienced slight distension of her abdomen as well as bouts of abdominal pain. In addition to this, she lost her appetite and suffered from fatigue. There had been no significant weight loss. Her medical history included an abdominal gunshot trauma sustained 4 years previously, which had resulted in laceration of the right liver lobe. The liver had been partially resected and a cholecystectomy was performed. There was no history of excessive alcohol intake or hepatitis.

At physical examination, we saw a pale woman with blood pressure of 110/65 mmHg. The abdomen was distended, with visible enlarged abdominal wall veins. A large firm mass was palpated in the right upper abdomen. There was pitting oedema and distension of the veins of both legs, consistent with an inferior vena cava syndrome.

Laboratory studies showed an erythrocyte sedimentation rate of 47 mm/h (10–20 mm/h), a leucytosis of 22.4×10\textsuperscript{9}/l (4–10×10\textsuperscript{9}/l), a haemoglobin concentration of 6.6 mmol/l (7.3–9.3 mmol/l), an alkaline phosphatase of 193 U/l (25–75 U/l), an alanine aminotransferase of 33 U/l (5–30 U/l), an aspartate aminotransferase of 58 U/l (5–30 U/l), a γ-glutamyl transpeptidase of 79 U/l (5–35 U/l), a lactate dehydrogenase of 730 U/l (160–320 U/l), a total bilirubin of 22 μmol/l (0.5–20 μmol/l), a creatinine of 155 μmol/l (60–110 μmol/l) and a urea concentration of 2.3 mmol/l (2.5–8 μmol/l). The urine sediment was normal. Computed tomography (CT) revealed, on transversal images, a large mass located in the

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right abdomen, possibly originating from the right kidney. An inferior vena cavogram showed complete occlusion of the inferior vena cava. Placing of a stent in the vena cava was technically not feasible. Magnetic resonance (MR) images were suggestive of an adrenal rather than a renal tumour. On coronal images a large tumour of about 25 cm in longitudinal axis was found (Fig. 1). These findings corresponded with the angiogram which revealed that the arterial blood supply of the tumour was derived from the right adrenal artery (Fig. 2). This arterial vascularisation produced evidence of the adrenal origin of the tumour.

A percutaneous thick-needle biopsy of the tumour had been performed in the referring hospital, and revealed a spindle cell tumour with features of smooth muscle differentiation; leiomyoma and leiomyosarcoma were considered in the differential diagnosis. CT and MRI showed no evidence of metastases to lungs, brain or lymph nodes.

An exploratory laparotomy was performed. A sub-hepatic tumour was found to extend from the midline to the right abdominal wall, with complete fixation to the liver. Because of this, resection was considered impossible, also in view of the poor clinical condition of the patient. Palliative and supportive care was instituted; the patient died 3 weeks after the laparotomy.

Autopsy revealed a predominantly sub-hepatic grey–white tumour with a maximal diameter of 27 cm. The right adrenal gland could not be identified. The left adrenal gland appeared to be normal. No tumour mass was found in the uterus or bowel or any other site at autopsy. Histological examination again revealed a spindle cell tumour with a moderate degree of atypia and with up to 10 mitoses per 2 mm², with areas of coagulation necrosis (Fig. 3a).

Tumour cells were elongated, had blunt-ended nuclei and the cytoplasm was eosinophilic with a slightly fibrillary appearance. Immunohistochemistry showed strong immunoreactivity for smooth muscle actin (Fig. 3b), attesting to the smooth muscle differentiation of the tumour. At autopsy, metastases or other primary sites were not identified. There was a right-sided pleural effusion resulting in atelectasis of the subjacent right lung. The vascular wall of the inferior vena cava was intact, excluding a possible primary tumour of the inferior vena cava. Thrombosis of the inferior caval vein was found, which could be explained by extrinsic compression by the tumour.

Discussion

Primary mesenchymal tumours of the adrenal gland are very rare. Most are benign, with myelolipomas and haemangiomas being the most common entities in this group. A literature search identified four previous reports of primary adrenal leiomyosarcoma. These occurred in three males and one female, ranging in age from 30 to 49 years. In about 200 cases described in literature a leiomyosarcoma originated from the inferior vena cava. These patients often presented with a inferior vena cava syndrome as well. Other primary sites of vascular leiomyosarcoma are even more rare.

In general, adrenal masses larger than 3 cm are suspicious of malignancy. To evaluate the resectability of a suspicious adrenal tumour, pre-operative imaging of the tumour and screening for distant metastases is essential. In view of the rarity of primary adrenal leiomyosarcoma, it is obvious that the possibility of a clinically occult tumour elsewhere needs to be carefully ruled out before this diagnosis is made. In our case CT and MRI could not detect any other sites of tumour growth. CT alone cannot provide definitive proof that a large upper abdominal tumour is of adrenal origin and angiography is of great value in the pre-operative work-up of such patients. But one must bear in mind that a metastasis in the adrenal gland would also be supplied by the adrenal artery. The combination of CT, MRI and angiography proved useful in this patient.

In our case, there is a previous history of a gunshot trauma necessitating a partial hepatectomy of the right liver lobe. In the literature there is a scattering of case reports that suggest that occasionally, severe tissue trauma such as may result from gunshots could be a causal or contributing factor in the pathogenesis of some sarcomas. At time of the emergency laparotomy for the lacerated liver, there had been no evidence of a tumour. Whether the adrenal gland had been damaged by the gunshot wound is not clear from the records. In our case the latency period between trauma and diagnosis of malignancy was 4 years; most cases of sarcoma arising after tissue trauma report a much longer period.
Histopathological evaluation is indispensable not only for determining tumour type but also for grading and, in parallel biological aggressive behaviour. In this respect the mitotic index and the presence of necrosis are important parameters. Lack et al. described a case of a high grade adrenal leiomyosarcoma with distant metastases at time of diagnosis which could be treated only palliatively. One of the other previously reported
cases of adrenal leiomyosarcoma concerned a low-grade tumour that showed no evidence of recurrence at 20 months after operation. In this fifth reported case of a primary leiomyosarcoma of the adrenal gland the patient did not die because of malignant behaviour of the tumour but because of the mechanical problems caused by this very large tumour. The patient suffered from a vena cava inferior syndrome, atelectasis of the right lung with 2.5-l pleural fluid and bronchopneumonia of the left lung.

Fig. 3. (A) Histological appearance of the tumour. A proliferation of elongated tumour cells with predominantly blunt-ended nuclei is seen. A mitosis is present left of centre (→). Haematoxylin and eosin, magnification, ×700. (B) The tumour cells are strongly positive (→) for smooth muscle actin, a protein involved in smooth muscle cell contraction and a histopathological marker of smooth muscle cell differentiation. Avidin–biotin peroxidase immunostaining, magnification, ×450.
It seems that aggressive local surgical management of low-grade sarcoma can produce prolonged survival in these patients. Because a radical resection or compartmental resection with wide tumour-free margins in the retroperitoneal space is not possible for large tumours, the local recurrence is high. From soft tissue sarcomas of the extremities it is known that adjuvant radiotherapy after a marginal resection results in similar survival rates as after a radical resection only. Therefore radiotherapy should always be considered after a marginal resection of retroperitoneal soft tissue sarcoma. The role of adjuvant chemotherapy in soft tissue sarcoma still remains unclear.

A meta-analysis of 14 randomised studies suggested a possible survival improvement, however not significant.

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