Epic battles in endocrinology — malignant pheochromocytoma: a case report

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Abstract
The rarity of malignant pheochromocytoma coupled with the lack of definitive predictors of malignancy and thevariability of clinical course, poses a significant diagnostic and therapeutic challenge. Since data on treatment is so scarce, case reports are a valuable source of knowledge for clinicians.

This case report describes the medical history of a woman, aged 51 at the time of initial diagnosis and adrenalectomy. Within over 5 years she presented with recurrent relapse of tumour in adrenal gland bed and multiple distant metastases to descending colon, abdominal wall, postoperative scars, and the peritoneum. Neither before diagnosis nor during the whole follow-up were symptoms associated with pheochromocytoma present.

The treatment administered to our patient consisted of numerous debulking surgeries along with administration of both hot and cold somatostatin analogues. We believe that debulking surgeries played a substantial role in enabling the patient to survive nearly 6 years despite aggressive clinical course of pheochromocytoma. She passed away in 2012 as a result of postlaparotomy complications.

We stress the role of debulking surgery in the treatment of malignant pheochromocytoma and summarise current literature.

Key words: malignant pheochromocytoma; debulking surgery; case report; lanreotide

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Introduction
Pheochromocytoma (pheo) is a rare catecholamine-secreting tumour, although it may also be nonfunctional [1]. Its prevalence is less than 5 per 1 million of the general population and less than 1% of the hypertensive population, with no significant gender predominance. Although in most cases benign, hormonally active pheo poses a real life threat due to the sympathetic system overstimulation. Malignant pheo constitutes 10–25% of all cases [2].

Given the rarity of this condition, therapeutic options are not well defined and those which are available are mostly palliative [3].

Case report
We present a case of a Caucasian female diagnosed with malignant pheo, treated with surgery along with both cold and hot somatostatin analogues. In October 2006, at the age of 51, she underwent an
Abdominal ultrasound (US) scan due to recurring pains of the lumbar area. A 109 x 89 x 92 mm mass of the left adrenal gland was visualized. Irregular echogenicity and focal necrosis were described (Fig. 1). Typical pheo-associated symptoms, such as paroxysmal hypertension, tachycardia, sweating, tremors or anxiety were not present at any time. Multiple blood pressure readings in 2006 during a hospitalization ranged up to 120–130/80 mm Hg.

A computed tomography (CT) scan performed in October 2006 confirmed the presence of a 90 x 95 x 120 mm lesion in the left adrenal gland with focal necrosis and irregular attenuation coefficient along with moderate enhancement after contrast administration. The edges were well defined and the tumour did not involve the adjoining tissues (pictures not available). Laboratory testing showed normal function of the adrenal cortex and medulla. Total metanephrine level in a 24-h urine sample was 300 µg (reference ranges < 1000 µg/24 h). 24-h fractionated metanephrines were not assessed since such testing was unavailable at our centre at the time.

A classic adrenalectomy was performed in November 2006. The excised tumour measured 140 x 120 x 90 mm and weighed 430 grams. Pathologic evaluation indicated pheo with no invasion beyond the capsule. Immunohistochemical staining was positive for chromogranin A and synaptophysin.

A CT scan performed in January 2007 showed no signs of recurrence; however, shortly after that the patient reported two tumours she palpated in the scar. In US two solid lesions in subcutaneous tissue were described. In March 2007 the nodules were excised and the histopathological examination indicated pheo. Diagnosis of malignant pheo was suspected. Bone scintigraphy performed in February 2008 revealed no pathological uptake. A routine, follow-up CT scan the patient underwent in May 2008. Examination revealed a 25 x 29 x 36 mm mass in the left adrenal gland bed together with 2 tumours in the scar area and descending colon wall thickening with considerable narrowing. Hormonal evaluation revealed no catecholamine excess again. Genetic testing for RET, VHL, NF1, SDHB, SDHC and SDHD mutations was negative.

Because of those findings our patient underwent a laparotomy in June 2008, when a left hemicolectomy was performed and the abdominal wall tumours were removed. The mass in the left adrenal bed was deemed inoperable due to infiltration of both superior mesenteric vein and left renal vessels.

At that time, the diagnosis was unclear. The clinical picture could indicate both pheo-paraganglioma syndrome and malignant pheo due to the fact that the masses appeared in the area that was previously operated on. However, malignant pheo was highly suspected. In order to investigate the therapeutic options the patient underwent a ¹²³I-MIBI scintigraphy along with ⁹⁹mTc-HYNIC-TOC and NeoSPECT ⁹⁹mTc-HYNIC-TOC scans, the results of which in-

Figure 1. An ultrasound picture of a large (109 x 89 x 92 mm) adrenal tumour
Dicated that somatostatin analogues would be the best option, with lanreotide as a first choice. $^{123}$I-MIBI was decided against due to insufficient tracer uptake.

In September 2008 the patient presented with abdominal pain and underwent a CT scan. A large mass $10 \times 9 \times 6$ cm in the right mid abdomen was revealed and a month later it was excised. Histopathologically, pheo was described and final diagnosis of malignant pheo was established (Fig. 2).

Follow-up CT (December 2008) revealed two pathological masses in primary tumour bed. The larger one was $42 \times 14$ mm and the smaller one was $19 \times 12$ mm. Radiolabelled lanreotide (Y90-Lan) was administered in November 2008 and January 2009. In February 2009 the patient underwent a surgical excision of a new pathological mass in the skin of left anterior superior iliac spine area.

A month later she presented with pancytopenia which caused the diagnosis of multiple myeloma to be considered. Full diagnostic workup was performed and multiple myeloma was excluded. Pancytopenia was attributed to hot somatostatin analogue treatment, which was therefore discontinued. Because of abdominal pain, another abdominal CT was performed. Examination revealed progression of previously described tumours and new pathological mass in pelvis (left ovary area) measuring $38 \times 31$ mm (CT scans not available).

In June 2009, after a consultation we decided on debulking surgery. Extensive procedure with tumour in left adrenal bed resection (in June 2008 described as nonoperative), left nephrectomy and antero-superior rectal resection and partial small bowel resection was performed. Histopathologically, every resected tumour contained pheo tissue.

Routine abdominal US revealed new pathological masses in skin and subcutaneous tissue. In May and September 2010 our patient underwent two surgical procedures. First, lesions in the abdominal wall were excised, and the second operation involved the removal of 2 retroperitoneal tumours. “Cold analogue” lanreotide therapy was started in September 2010 and was continued throughout the whole follow-up period.

In February 2011 a positron emission tomography (PET) scan showed high risk of peritonitis carcinomatosa and therefore, in April 2011, a laparotomy was performed. At the time an intramural tumour of the rectum was removed together with 26 tumours from the retroperitoneum.

In June 2011 pathological masses behind the front abdominal wall were discovered. No surgical procedure was performed. In March 2012 the patient presented with nausea, abdominal pain together with rapid increase in creatinine levels and oliguria, which led to the diagnosis of hydronephrosis. A double J stent was placed in the right ureter which alleviated the patient’s symptoms. A CT scan performed at that time revealed further disease progression with large tumour in primary tumour bed ($85 \times 65 \times 100$ mm), few peritoneal implants and new 17 mm pathological mass in pelvis compressing right ureter. The patient underwent a laparotomy. The perioperative period was complicated with a severe haemorrhage, which was fatal. Prior to the last laparotomy the patient was offered treatment with chemotherapy, which she declined. She had

Figure 2. A CT scan of a large pathological mass ($10 \times 9 \times 6$ cm) in the right mid abdomen
never had typical symptoms of pheo. All procedures are summarized in Table 1.

**Discussion**

Due to rarity and variability of clinical presentation, pheo is a real diagnostic challenge for physicians [2]. The clinical course may be highly variable and no definitive predictive factors of malignancy are currently available. Recent data shows that tumours with *SDHB* mutation present significantly higher rates of malignancy [4]. Owing to the fact that a remarkable number of patients still die due to hormone overproduction, the main treatment objectives are controlling the tumour mass and hormonal symptoms [3]. Pheo is considered a secondary cause of hypertension, one that can be corrected surgically. Therefore it needs to be taken into account in every patient presenting with an adrenal tumour together with hypertension (paroxysmal or persistent). Even in the absence of typical symptoms and elevation of biochemical markers one cannot exclude pheo (malignant or benign) [5].

The diagnosis of malignancy can only be made by the discovery of distant metastases, direct invasion of surrounding tissues or presence of pheo in sites where chromaffin tissue is normally absent [6]. Pathologic markers are not predictive enough to be considered reliable in terms of tumour malignancy [2, 3]. Most common metastatic sites include liver, lungs, lymph nodes and bones [7]. Metastases may appear as long as 20 years after initial presentation, therefore long-term follow-up is mandatory [8].

Data concerning survival rates varies — some patients present a more aggressive clinical course and die within 5 years of the diagnosis, whereas others show no disease progression for many years despite not receiving any targeted therapy [2]. In the literature we reviewed, survival times ranged from 1 year to 23 years [2, 9]. However, since there are no reliable prognostic factors, determining the aggressiveness of the clinical course is impossible.
Therefore a dilemma naturally arises — whether to treat and when to treat.

Given the rarity of malignant pheo, there definitely is a lack of prospective and randomized studies. Available data is not completely reliable as reports on efficacy of different treatment modalities are variable. Having analysed a number of reports, we feel that the following findings are worth mentioning.

Chemotherapy seems to bring the most benefit to patients presenting with rapidly progressive disease rather than those with a more indolent clinical course [3], although Tanabe et al. found it to be more effective in the latter group [10]. The use of CVD (cyclophosphamide, vincristine, dacarbazine) regimen is well-recognized. Its implementation was justified by the resemblance between chromaffin tumours and neuroblastomas [11]. Adverse effects are mostly mild, the most common being gastrointestinal symptoms, leukopenia and hepatic dysfunction. If there is no significant response (biochemical or tumoural) after 4–5 cycles of CVD, it should be discontinued [10]. For inoperable lesions or multiple hepatic metastases, transcatheter arterial embolization with concomitant administration of doxorubicin or mitomycin-C has been attempted [12, 13].

In the case of malignant pheo surgery should be considered as a treatment of choice for both primary tumours and metastases [14]. Its intent may be to cure the disease or alleviate hormonal as well as compression-related symptoms [15]. Moreover, debulking surgery may also improve response to later treatment [8, 15]. Prior to surgery, as for every other treatment in the case of a hormonally active disease, adequate alpha-adrenergic blockade is mandatory [16, 17]. This prevents intraoperative complications related to massive catecholamine release, such as a hypertensive crisis, arrhythmias and myocardial infarction. Although surgical resection is the only potentially curative approach, it is rarely feasible due to delayed diagnosis of malignancy. In some cases metastasectomy proved to be a sufficient treatment, leading to complete remission and patients remaining asymptomatic for years [18, 19]. Our patient underwent multiple debulking surgeries which, together with long-term lanreotide treatment, enabled the almost six-year follow-up that she achieved. Reducing tumour burden might have facilitated better outcome of lanreotide therapy since cold somatostatin analogues alone have not shown considerable efficacy in treatment of malignant pheo.

Approximately 36% of patients diagnosed with malignant pheo are carriers of a hereditary germline mutation of SDHB (succinate dehydrogenase sub-unit B) [20]. Malignant pheos, particularly those associated with SHDB mutation, have been reported to be highly vascularized tumours. Therefore, anti-angiogenic therapy with sunitinib was implemented. Although not curative, it can be an alternative for patients with progressive disease resistant to therapy or not eligible for other treatment modalities [20]. The effectiveness seems to be satisfying, as sunitinib was reported to lead to at least disease stabilization. Shen et al. described necrosis of majority of the metastatic masses as a result of this course of treatment [21]. Side effects included hypothyroidism, hypertension and haematological toxicity [9, 21, 22].

123I-MIBG treatment has been attempted in various dose settings. According to Rose et al., repeated high-dose 123I-MIBG may be associated with long-term survival [23]. Single high-dose administration was also proven effective. However, this kind of treatment may result in a high percentage of severe haematological toxicity. Multiple intermediate or low doses regimens were more tolerable and had a reduced rate of myelosuppression while still offering a chance of prolonged survival [24–27]. Even in the case of “non-responders” in terms of tumour shrinkage or biochemical parameters, this course of therapy brought considerable symptomatic relief, therefore improving the patients’ quality of life. In order to prevent hypothyroidism, thyroid blockade should be attempted even though it has been reported that despite potassium iodate use, tracer uptake to the thyroid is still visible on the scans [25].

If the lesions show sufficient tracer uptake on somatostatin receptor scintigraphy, treatment with somatostatin analogues may be attempted [27]. Generally, radioactive analogues are recommended, as “cold” ones have resulted in only transient responses. Lamarre-Cliche et al. proved that slow-release octreotide administered in the group of patients who did not undergo debulking surgery or receive other treatment such as MIBG or embolization appears to not be of sufficient effectiveness for the long-term treatment [28]. Moreover, it does not improve pheo-related symptoms significantly. Side effects of radioactive SST analogues are mainly haematological, as shown in the case of our patient, who became pancytopenic due to treatment with a hot SST analogue and had to be treated with non-labelled lanreotide instead.

**Conclusion**

The rarity of malignant pheochromocytoma, lack of definitive predictors of malignancy and the vari-
ability of clinical course pose a significant diagnostic and therapeutic challenge. Since data on treatment is so scarce, case reports are a valuable source of knowledge for clinicians.

Debulking surgery may play a substantial role in prolonging the patients' survival.

References


