Castration failure in prostate carcinoma due to a functioning adrenocortical carcinoma

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Summary

We report concurrent metastatic prostatic adenocarcinoma (PC) and functioning androgen-secreting adrenocortical carcinoma (ACC) in a 77-year-old man. The failure to achieve adequate biochemical castration via androgen deprivation therapy (ADT) as treatment for PC metastases, together with elevated DHEA-S, androstenedione, and discordant adrenal tracer uptake on FDG-PET and PSMA-PET, suggested the presence of a concurrent functional primary adrenal malignancy. On histopathological analysis, scant foci of PC were present throughout the ACC specimen. Castration was achieved post adrenalectomy with concurrent drop in prostate-specific antigen. We outline the literature regarding failure of testosterone suppression on ADT and salient points regarding diagnostic workup of functioning adrenal malignancies.

Learning points:

• Failure to achieve castration with androgen deprivation therapy is rare and should prompt careful review to identify the underlying cause.
• All adrenal lesions should be evaluated for hormone production, as well as assessed for risk of malignancy (either primary or secondary).
• Adrenocortical carcinomas are commonly functional, and can secrete steroid hormones or their precursors (androgens, progestogens, glucocorticoids and mineralocorticoids).
• In this case, a co-incident, androgen-producing adrenocortical carcinoma was the cause of failure of testosterone suppression from androgen deprivation therapy as treatment for metastatic prostate cancer. Pathological adrenal androgen production contributed to the progression of prostate cancer.

Background

The primacy of androgen signalling in the development of prostate cancer is well defined (1). Androgen deprivation therapy (ADT) has been the cornerstone of metastatic prostatic adenocarcinoma (mPC) management for decades, with prolonged survival achieved via pharmacological therapies to lower circulating testosterone (2). The majority of advanced prostate cancer patients respond initially to ADT, with a target castrate level defined as testosterone <0.7 nmol/L. Failure to achieve a castrate level of testosterone is rare, with only isolated case reports in the literature (3). Here we present the first reported case of failure of testosterone suppression despite medical ADT in a patient with mPC due to a concurrent androgen-secreting adrenocortical carcinoma.
**Case presentation**

A 77-year-old man presented with a 12-month history of worsening mid-thoracic back pain. Initial workup revealed elevation of serum prostate-specific antigen (PSA) at 490 µg/L (reference range (RR): <7.5). A transrectal prostate biopsy confirmed Gleason 5+4=9 prostatic adenocarcinoma (PC). Staging contrast-enhanced computed tomography (CT) showed an enlarged prostate, widespread sclerotic bony lesions, malignant retroperitoneal lymphadenopathy and visceral lung lesions. Additionally, the left adrenal gland was enlarged by a heterogenous, 68 x 45 x 45 mm mass without calcification but with elevated pre-contrast density (20–30 Hounsfield units), with significant contrast uptake and slow washout (Fig. 1E and F). A clinical diagnosis of metastatic prostate cancer was made, with the identified lesions ascribed to prostate cancer metastases, and he was commenced on ADT with leuprorelin (GnRH analogue) and cyproterone acetate (androgen receptor antagonist).

Following 8 weeks of ADT therapy, PSA had fallen to 120 µg/L; however, serum testosterone remained elevated at 21.5 nmol/L (reference range 8–30), despite undetectable

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**Figure 1**

(A and B) Pre-operative 18F-FDG-PET, cine (A) and axial-fused (B), demonstrating focal intense uptake in left adrenal gland (blue arrow), with minimal uptake in the skeleton body (green arrows). (C and D) Pre-operative 68Ga-PSMA-PET, cine (C) and axial-fused (D), demonstrating modest uptake in left adrenal gland (blue arrow) but intense widespread uptake in the skeleton and vertebral body (green arrows), consistent with metastatic prostatic carcinoma. (E and F) Pre-operative axial computed tomography without intravenous contrast (E) and with contrast in the arterial phase (F), demonstrating a large, irregular, heterogeneous left adrenal gland with pre-contrast density 20–30 Hounsefield units. The right adrenal gland can be seen as normal. (G and H) 12 months post-operative 18F-FDG-PET, cine (G) and axial-fused (H) showing a new, intensely hypermetabolic liver metastases (gold arrow). Thoracic and abdominal nodal metastases are also seen (G).
levels of follicular stimulating hormone (FSH) and luteinising hormone (LH) (Table 1). Given persistently elevated testosterone, the GnRH analogue was switched to triptorelin. Despite this, testosterone remained persistently elevated at 23 nmol/L, with PSA 98 µg/L. Repeat staging CT showed growth in the left adrenal mass (now 78 x 50 mm) with local extension towards the posterior surface of the stomach. Other sites of metastatic disease were stable.

**Investigation**

The differential diagnosis for the enlarging left adrenal mass was reviewed. Clinical examination revealed no evidence of androgen or cortisol excess. He appeared eugonadal, uncommon on ADT. An adrenal hormonal panel showed, in addition to persistently elevated testosterone performed in two separate assays, elevated DHEA-S at 13 µmol/L (RR: <10) and androstenedione 28 nmol/L (RR: 2.1–11). Evidence for autonomous cortisol excretion was equivocal with normal 24 h urinary free cortisol (159 nmol/24 h, RR: <270) but failure of suppression of cortisol following 1 mg overnight dexamethasone suppression test (basal cortisol 437 nmol/L with ACTH 0.6 nmol/L (RR: 1–10); post-suppression cortisol 363 nmol/L, normal: <50 nmol/L). Plasma metanephrines and normetanephrines were negative. The presumptive diagnosis for the left adrenal lesion was revised to androgen-secreting adrenocortical carcinoma (ACC).

Functional imaging with fluorine-18 fluorodeoxyglucose (FDG) positron emission tomography (PET) scan identified intense radiopharmaceutical uptake at sites of skeletal metastases (Fig. 1A and B). By contrast, a Gallium-68 prostate-specific membrane antigen (PSMA) PET showed moderate radiopharmaceutical uptake within the ACC and avid radiopharmaceutical uptake within the metastases throughout the skeleton (Fig. 1C and D). This, combined with the structural stability of the skeletal lesions, was considered supportive that the metastatic disease was prostatic carcinoma and that the presumed ACC remained localised to the adrenal bed.

**Treatment**

In view of the high lethality of untreated ACC, as well as the strong clinical suspicion that androgen secretion from the adrenal tumour was driving progression of metastatic prostatic cancer, he proceeded to left adrenalectomy. Given the presence of prostatic cancer metastases, a laparoscopic approach was attempted to minimise morbidity. However, the tumour was highly vascular and difficult to mobilise, requiring conversion to an open procedure. The adrenal tumour was removed intact with marginal clearance but preservation of all surrounding structures. There were no perioperative complications.

Histological examination showed a pT3 high-grade adrenocortical carcinoma (Ki-67: 40%), 80 mm in maximal length, invading perivesical fat but with clear resection margins. Scant foci of metastatic PC were observed within the ACC parenchyma (Fig. 2). These concomitant carcinomas were confirmed on immunohistochemical staining, with mutually exclusive synaptophysin (Fig. 2C) and inhibin expression in ACC (Fig. 2C) and prostate-specific antigen expression in mPC (Fig. 2D).

| Table 1 | Hormonal profile prior to and following treatment for adrenocortical carcinoma. |
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| Event | Pre-adrenalectomy, day | Post-adrenalectomy, day | Reference range |
| | D-120 | D-90 | On ADT | ADT + Mitotane | ADT + Mitotane | ADT + Mitotane | Mitotane ceased | Palliative pathway |
| PSA, µg/L | 490 | 98–120 | 16.0 | 0.20 | 0.92 | 5.4 | 4.7 |
| Testosterone, µmol/L | 21–23 | <0.4 | <0.4 | <0.4 | 7.3 | 16.9 |
| DHEA-S, µmol/L | 10–13 | 0.9 | 0.1 | 0.6 | 15.8 | 25.4 |
| Androstenedione, nmol/L | 28 | 0.9 | 0.9 | NA | 32.1 | 55.0 |
| ACTH, pmol/L | <0.1 | 1.0–10.8 |
| Cortisol, nmol/L | 437 | 91 |
| 8 am, basal post 1 mg DST | 363 |

Day (D) is with reference to day of adrenalectomy Day = 0; ADT, androgen deprivation therapy; DST, cortisol level following 1 mg overnight dexamethasone suppression test; PSA, prostate-specific antigen.
Outcome and follow-up

Three weeks post-operatively PSA fell to 30 µg/L, with a now appropriately suppressed testosterone at <0.4 nmol/L (Table 1). Six weeks post-operative PSA was 16 µg/L with low DHEA-S 0.9 µmol/L, androstenedione 0.9 nmol/L and testosterone <0.4 nmol/L. He received adjuvant irradiation to his adrenal bed. Given the limited prognosis even in resected stage III ACC, with a 24% 5-year survival only, the consensus multidisciplinary team decision was to prioritise adjuvant treatment for his ACC over intensifying systemic therapy for his mPC, and adjuvant mitotane was commenced. Standard of care GnRH-based ADT for mPC, now in the metastatic hormone-sensitive state (mHSPC), was continued. The patient remained stable on mitotane and ADT, in clinical and biochemical remission from ACC, for 12 months after adrenalectomy. A surveillance 18F-FDG PET at 6 months showed no evidence of recurrent disease. However, 18F-FDGPET performed 13 months after adrenalectomy, and coincident with the development of thoracic back pain and deconditioning, demonstrated multiple FDG-avid foci consistent with metastatic ACC (Fig. 1G and H). Concurrently, levels of adrenal androgens began to increase, consistent with functional ACC metastases (Table 1), with corresponding increases in PSA, consistent with stimulation of mPC by adrenal androgens. Due to a declining functional status, palliative chemotherapy was not initiated. Mitotane therapy was ceased, and palliative care was enhanced. He died 16 months after adrenalectomy and 20 months after diagnosis of metastatic prostate cancer.

Discussion

This case represents an unusual cause of failure of medical ADT due to a functioning, androgen-secreting, ACC. ADT, via medical or surgical approaches, has been a cornerstone of treatment for advanced prostate cancer since the first work outlining the sensitivity of this disease to androgens. In this case, adrenal androgens contributed to the progression of mPC, given the correlation between PSA and androgen status both at initial diagnosis and at ACC recurrence.

Given gonadotrophin receptor agonists/antagonists are widely effective medical approaches to achieving castration, they are established first-line therapy for androgen deprivation in advanced prostate cancer. There are isolated case reports of primary failure of suppression of testosterone secretion in this setting (4), with various possible explanations offered, ranging from anti-drug antibodies, incorrect administration or rapid metabolism of drug. Persistent elevation of LH suggests incorrect administration. A single prospective observational study reported that the rate of failure of ADT to suppress testosterone was 5% (n = 38); associated with obesity, and resolved following orchidectomy (5). A single case of a functional gonadotrophin-secreting pituitary adenoma has been reported (6).

Adrenocortical carcinomas (ACC) are rare, with a bimodal age distribution affecting young children and those in middle age, with a slight female preponderance. Sixty per cent are sufficiently secretory to present with symptoms of hormone excess, such as Cushing’s
surgical resection of the ACC may not only improve outcome for the prostate cancer (by reducing androgen secretion) but also may improve outcome from the primary adrenal malignancy by resecting all known disease.

Crowley and colleagues also postulated the potential of using lutetium and actinium PSMA labelled radioisotopes as a treatment option for ACC (9). In this case where there is a synchronous prostate cancer, PSMA radionuclide therapy would have the advantage of potentially working for both the distant metastatic prostate cancer and the future recurrent or metastatic ACC.

Mitotane, an insecticide isomer, is an adrenolytic drug used to treat ACC. It has a cytotoxic effect on mitochondria in the adrenocortical cells of the reticularis and fasciculate. Mitotane inhibits activity in the cholesterol side chain cleavage (CYP11A1) enzyme and CYP11B activity within ACC, affecting adrenal cortisol and testosterone biosynthesis. Indeed, when used in the adjuvant setting for ACC, mitotane has been found to decrease DHEA-S concurrent with cortisol, whilst being associated with a clinical syndrome of hypogonadism. Extra-adrenal actions of mitotane and increased sex hormone-binding globulin (SHBG) and resultant decreased free testosterone, together with inhibition of 5α-reductase, contribute to known hypogonadal sequelae. A human hepatoma cell line model utilising male patients on adjuvant mitotane for ACC suggested reduced total and free [testosterone], concurrent with oestrogen- dependent increase in SHBG, whilst being associated with a hypogonadal state induced by the pluripotent activity of mitotane. This hypogonadal state induced by the pluripotent activity of mitotane may provide a beneficial impact on the clinical course of our patient’s castrate-sensitive PC, independently of medical ADT and resection of his androgen-producing ACC. Murine mHRPC xenograft studies have shown that adrenally derived androgens are key drivers of tumour growth, with surgical bilateral adrenalectomy reducing tumour volumes (10).

This unusual presentation of a concurrent functioning ACC and mPC illustrates the complex endocrine interplay of both diseases. The failure of castration via ADT is a rare occurrence and should prompt careful evaluation of a wide range of differential diagnoses.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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