Metastatic Cancer of Cowper’s Gland: A Rare Cancer Managed Successfully by Molecular Profiling

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Abstract
Cancer of Cowper’s gland is a very rare cancer. This case represents the 9th case in the medical literature. As such, there are no phase II or phase III trials to guide treatment. In this article, we report the successful treatment of a patient over a 7-year period guided solely by molecular profiling. Through multiple cycles to treatment, the cancer was controlled using drugs targeting c-kit, as the cancer steadily increased the expression of c-kit. This report also documents the use of a novel drug combination based on sunitinib that was well tolerated and may warrant testing in other c-kit-dependent cancers.

Introduction
Cancer of Cowper’s gland is a rare cancer with fewer than 10 recorded cases [1–6]. The literature is composed of case reports, primarily predating the modern era of chemotherapy. This means making treatment decisions without randomized phase III or even phase II trials.

The reported cases of Cowper’s gland carcinoma have all been adenoid cystic carcinoma (AdCC), and treatment effective against AdCC arising in other organs might be used. However, AdCC is uncommon or rare in each of the organs in which it is found. This pathology is most commonly found in salivary gland cancers, which constitute less than 5%
of all head and neck cancers. As a result, there are no randomized phase III clinical trials with sufficient statistical power to guide management of AdCC at any organ site. Even phase II clinical trials are limited in number, and none has yielded impressive therapeutic results. Most clinical studies in AdCC show only disease stabilization. Objective tumor responses are uncommon [7–16]. Furthermore, the available information suggests there may be significant differences in drug effectiveness in AdCC arising in different organ sites [17–19].

An alternate approach to uncommon cancers is to let molecular profiling guide selection of drugs for systemic therapy. The attraction of this approach is that the treatment is selected based on known mechanisms of drug action. Furthermore, accumulation of such information is likely to shed light on how AdCC at various organ sites differs in ways that can be used not only to guide therapy of individual patients, but also to guide clinical trial design. In this article, we report the results of this approach with a case of Cowper’s gland AdCC.

Case Report

The patient initially presented with pelvic pain and rectal discomfort. CT showed a prostate mass. Based on a core biopsy, he was diagnosed with Cowper’s gland AdCC.

When the patient initially presented for treatment, we faced the quandary posed by the lack of clinical trial data on this very rare disease. The options were no treatment at all or to let molecular profiling guide treatment choices. This issue was discussed in detail with the patient and his wife, and they elected to proceed with treatment. The case and the treatment quandary were presented to the hospital tumor board, and the decision to start treatment was agreed upon. At every step in the course of the treatment outlined, decisions were solely based on the treatment options that seemed to offer the patient the best balance between cancer control and quality of life.

The patient was initially treated with pelvic exenteration on February 26, 2003. However, the surgical margins were positive, and he received adjuvant radiation, cisplatin and taxol. He remained disease free for just under 3 years, until he experienced recurrence on January 11, 2006, with a mass in the surgical bed. A biopsy confirmed recurrent AdCC. Immunohistochemistry documented the presence of c-kit and epidermal growth factor receptor (EGFR). No treatment was offered. By July 2006, the patient had documented liver metastasis seen on CT, and these lesions were biopsy positive for AdCC. The mass in the surgical bed was removed and the liver lesion was treated with cryoablation.

While AdCC can often progress slowly, this patient demonstrated rapid growth and spread of his disease. By September 2006, he had developed a 1.3 × 1.3-cm lung lesion and an additional 2 × 2-cm liver lesion (fig. 1a). In October 2006, based on the initial immunohistochemistry detection of c-kit, the patient was started on a sunitinib-based combination (50 mg daily) because of its activity against c-kit as well as its impact on vascular endothelial growth factor (VEGF) and platelet-derived growth factor (PDGF) receptor kinase activity. The histone deacetylase (HDAC) inhibitor, valproic acid 500 mg, was added twice a day; thalidomide 50 mg and sargramostim 250 μg were added daily. The rationale for this combination is discussed below.

By December of 2006, six weeks after initiation of therapy, the lung lesion had disappeared, but the liver lesion remained stable (fig. 1b). Three months later (March 2007), the liver lesion had increased marginally to 2.5 × 2.0 cm. At this point, the liver lesion was treated with cryoablation. At the end of May 2009, the patient had a screening colonoscopy with perforation of the bowel, leading to sepsis and suspension of cancer treatment for almost 1 month. During this time, he developed a rapidly progressing lesion at L4. Thus, the sunitinib-
based combination resulted in a progression-free survival of almost 2.5 years, during which he was able to work fulltime as a dentist. The sunitinib-based combination was restarted but proved ineffective. Because of EGFR expression, he was treated in turn with erlotinib and lapatinib without clinical benefit but with progression of pulmonary metastases.

At this point, the surgical specimen from May 2009 (metastatic carcinoma from T6 epidural space) was submitted to Caris Life Sciences, Phoenix, Ariz., USA, for more detailed analysis (Target Now tumor profiling). This specimen showed overexpression of TOPO1 (fig. 2a), suggesting sensitivity to irinotecan. As a result, he was started on cycles of irinotecan 125 mg/m² per week × 4, followed by 2 weeks off. His disease remained stable for 10 months before he progressed.

The Caris Target Now analysis of the July 2010 sacral biopsy specimen had also shown continued expression of c-kit (fig. 2b). For this reason, in September 2010 he was started on imatinib 400 mg per day. He showed stable disease on imatinib for 9 months until he progressed with bone lesions leading to cord compression, which was managed with surgical resection.

In November 2011, he was started on gemcitabine based on overexpression of mRNA for deoxyctydine kinase. He did not respond, perhaps due to overexpression of RRM1, which has been linked to gemcitabine resistance [20]. In February 2012, he was started on liposomal doxorubicin because of overexpression of TOPO2B on RNA microarray. He derived substantial pain relief, with pain levels going from 7–8 to 2–3/10. The palpable mass arising from the thoracic spine showed marked reduction in size, without healing in the corresponding bone. In contrast, his renal lesions progressed on treatment. Doxil was terminated in August 2012.

Molecular profiling of a spinal (T6 epidural) tumor performed in August 2012 revealed overexpression of SRC mRNA, and the patient has been receiving dasatinib with good cancer control. He had an isolated site of progression in the spine that was successfully debulked surgically, after which dasatinib was continued. Additionally, molecular profiling of kidney metastasis of the AdCC revealed no mutations in KRAS, PIK3CA and BRAF genes.

**Discussion**

AdCCs of the salivary glands and breast have been associated with the presence of MYB-NFIB fusion, resulting in overexpression of MYB mRNA transcript, which was observed in our case as well (67 times the normal salivary gland control using Illumina microarray methodology) [21–25]. Pathologically, these cancers are composed of a dual cell population of luminal and myoepithelial cells. These cancers are often c-kit positive, but c-kit expression is commonly limited to the luminal cell component [21, 26, 27]. Additionally, c-kit is only rarely activated through gene mutations in AdCC, and no mutations were observed in our case [28]. The 5’ end of the c-kit gene contains MYB response elements, and MYB has been shown to up-regulate c-kit expression. Despite these findings, the role of c-kit (and possibly its ligand) remains to be established at a molecular level in the biology of these cancers.

By the fall of 2006 when we first encountered this patient, known treatment options for Cowper’s gland carcinoma had been exhausted. Furthermore, over a 3-month period, he had shown rapid cancer progression in the lungs and liver, suggesting a relatively poor prognosis. The presence of c-kit overexpression was the only clue available upon which to design a treatment program, and we chose to target this protein. Subsequent analysis showed overexpression of PDGF-C and PDGFRA at the RNA level and PDGF receptor by immunohistochemistry.
For his initial treatment program, we selected sunitinib because of its activity against c-kit [29, 30]. We elected to combine this drug with agents that might amplify the actions of sunitinib. At the time the combination was assembled, it was known that sunitinib reduced tumor-induced immunosuppression. Subsequently, sunitinib has been shown to diminish myeloid-derived suppressor cells and regulatory T cells [31, 32]. We elected to add low-dose thalidomide and sargramostim to further enhance any beneficial impact sunitinib might have on immune function.

Sunitinib also blocks angiogenesis via its capacity to block VEGF receptor and PDGF receptor, and his tumor expressed both PDGF receptor and its ligand. In this combination, valproic acid may have contributed because it has been reported to reduce expression of both VEGF receptor and VEGF. Additionally, valproic acid has been reported to be selectively toxic to c-kit+ GIST compared to c-kit− GIST [33].

This combination was very well tolerated and allowed the patient to continue to work at his chosen profession of dentistry. Additionally, it resulted in a partial response that lasted approximately 2.5 years. This combination may warrant additional testing in c-kit-positive malignancies.

Subsequent to this initial response, he has had disease stabilization and relief of pain from irinotecan, imatinib and liposomal doxorubicin, but no partial or complete remissions. Each of these treatments was chosen based on molecular profiling. The patient benefited from 3 out of the 4 treatments based on molecular profiling performed on the May 2009 surgical specimen. We must not overlook the undeniable contribution of the multiple surgeries which provided both symptomatic relief and samples for profiling, but also contributed to the prolonged survival of the patient. Whether or not the profiling was simply associated with or actually caused the responses seen cannot be determined outside the context of a randomized controlled trial. However, continuous profiling of the disease provided therapeutic monitoring and therapy adjustments, and also showed remarkably subtle changes in monitored biomarkers during the evolution/drug selection pressure of the disease process.

Despite having a cancer that rapidly progressed off treatment, our patient remains alive 6 years after progressing through known treatment options for Cowper’s gland carcinoma. We conclude that for rare cancers such as this, treatment selection based on molecular profiling may well prove to be a reasonable option.

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Fig. 1. Status of the pulmonary metastases prior to the treatment with the sunitinib-based combination (a) and in December 2006 after a response to the treatment (b).

Fig. 2. a Positive IHC staining for TOP01 expression. b Positive IHC staining for c-kit.