Percutaneous sequential bacillus Calmette-Guèrin and mitomycin C for panurothelial carcinomatosis

Savino M. Di Stasi, MD, Antonella Giannantoni, MD, *Robert L. Stephen, MD, Luigi Storti, MD, Francesco Attisani, MD, Andrea De Carolis, MD, Guido Virgili, MD

Department of Surgery/Urology, “Tor Vergata” University, Rome; Department Of Urology, University of Perugia, Perugia; Physion Laboratories, Medolla; Italy


A 59 year old male presented with a 4 month history of lower urinary tract symptoms.

Exhaustive urological investigations revealed papillary tumors and carcinoma in situ extending from the prostatic urethra, throughout the bladder, up both ureters and into the renal pelves.

Tumors were resected where possible and then bacillus Calmette-Guèrin (BCG) and mitomycin C (MMC) were infused sequentially through bilateral nephrostomy tubes for a total of six BCG and three MMC instillations.

Follow up 1 month post treatment demonstrated a complete response which persisted for 2 years. Then there appeared a solitary papillomatous recurrence in the bladder which was successfully resected. Side effects were the occasional fever and BCG induced granulomatous prostatitis which slowly resolved.

In conclusion, sequential BCG/MMC instillations were effective treatment for widespread panurothelial carcinomatosis.

Key Words: panurothelial, carcinomatosis, immunotherapy, chemotherapy

Introduction

Although transitional cell carcinoma (TCC) confined to the bladder is the most common site for urothelial cancers,1 panurothelial TCC occurs in significant numbers of patients with varying combinations involving bladder, upper urinary tract (UUT) and prostatic urethra.2 Most reports describe these combinations as metachronous3 with varying time intervals between detection of carcinoma at the different sites.4 Synchronous presentation with carcinoma extending throughout the whole urinary tract is rare. French investigators have described one such probable case but, as the full extent of the disease was defined over a time span of 3 years, the authors themselves were uncertain as to the sequence of events.5

In this report we describe a patient who presented with panurothelial carcinomatosis extending from the prostatic urethra to the renal tract...
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Case report

Initial findings
A 59 year old man presented with a history of hematuria, urinary frequency and dysuria of 4 months duration. Further questioning elucidated progressive reduction of urinary flow, nocturia (1-2 times) and incomplete bladder emptying that had been present for 2 years.

The patient had always worked as a post office employee with no exposure to occupational carcinogens but had smoked 30 cigarettes daily since age 14 (45 years). There was nothing relevant in his past medical history, he was taking no medications, had no allergies and the family history was unhelpful. Physical examination revealed chronic bronchitis, emphysema and slight prostatic enlargement. A urine sample contained malignant cells. An abdominal ultrasound and intravenous pyelogram with micturating cystourethrogram showed one irregular defect in the bladder neck, trigone and posterior bladder wall and another in the prostatic urethra at the verum montanum.

Urethrocytosispycoscopy revealed multifocal papillary tumors in the prostatic urethra, Figure 1 panel A, and in the bladder corresponding to the regions of the filling defects. Papillary lesions surrounded and also emerged from both ureteric orifices, Figure 1 panels B and C. All visible tumors were resected (TURBT) and their predominant pathology was transitional cell carcinoma grade G2 penetrating the lamina propria (T1). Obviously there was a need (a) to define the full extent of the carcinomatosis and (b) to initiate appropriate therapy.

Further investigations
A spiral CT scan of the upper urinary tracts (UUT) and ureteropyelography were uninformative. Endoscopic evaluation involving lavage of both UUT for cytology, bilateral ureteropyeloscopy and multiple biopsies of the UUT, the bladder and prostatic urethra delineated the clinical situation. Both ureters contained multiple, flat, hyperemic plaques and some small papillary tumors extending from the bladder up into the renal pelves. Pathology of the tumors was Ta G2 and that of the other biopsies was carcinoma-in-situ (Tis) extending from the prostatic urethra, throughout the bladder and up into the renal pelves: panurothelial Tis with multiple papillary tumors. There was no evidence of cancerous progression into muscle layers and beyond.

Treatment plan
Alternatives: The patient resolved the contentious issue of surgical extirpation – combined bilateral nephro-uretectomy and cystoprostatectomy – by refusing this option. A proposed experimental treatment with Pasteur bacillus Calmette Guèrin (BCG) and mitomycin C (MMC) was approved by the institutional review board and the patient supplied written informed consent.

Procedures: With ultrasound and fluoroscopy control bilateral percutaneous nephrostomy tubes were inserted into the superior renal calyces under local anesthesia. Unobstructed flow from the renal pelves to the bladder was assured and pyelovenous/pyelolymphatic backflow were excluded with fluoroscopy. The bladder was catheterized, with thorough drainage achieved under ultrasound control, then the catheter was removed. The drug solutions prepared for instillations were BCG 54 mg in 100 ml 0.9% NaCl solution and MMC 40 mg in 100 ml water.

Figure 1. Endoscopic pictures of the prostatic urethra (panel A), right (panel B) and left (panel C) ureteric orifices were taken during the initial transurethral resection.
and were divided into two 50 ml aliquots attached to
the two nephrostomy tubes. An immediate bilateral
instillation of 10 ml-15 ml theoretically filling the UUT
was followed by slow infusion of the remaining drug
solution over 1 hour. All symptoms during treatment
and the patient’s subsequent overnight stay were
recorded. Antibiotic prophylaxis was not employed.

Treatment cycles: The three sequential BCG-MMC
cycles totaled six BCG and three MMC instillations
as shown in Figure 2.

Follow-up
Response to therapy was assessed 1 month after
the last treatment cycle employing: urinary cytology,
saline lavage of both UUT and the bladder
with cytology, urethrocystoscopy, bilateral
ureteropyeloscopy, and selected biopsies of the
renal pelves, ureters, bladder and prostatic urethra.

Urinary cytology, urethrocystoscopy, bladder
biopsies and saline lavage of UUT with cytology
were repeated at 3 monthly intervals. An excretory urogram
and ureteropyeloscopy were performed yearly.

Results

Response to treatment
The investigations undertaken 1 month post treatment
indicated a complete response. We found no evidence
of carcinoma along the whole length of the
urothelium. Both nephrostomy tubes were removed
and the response persisted for 2 years. Symptoms
indicative of minor bladder outlet obstruction were
relieved by Tamsulosin 0.4 mg daily and no other
problems appeared.

Recurrence
Two years after completion of the anti cancer
treatment, suspect tumor cells were detected in the
urine. Cystoscopy revealed a papillary tumor 5 mm
in diameter on the posterior bladder wall. Generous
resection (TURBT) and pathological examination
showed transitional cell carcinoma grade G2 stage T1.
The exhaustive investigations of the first follow up
examination were repeated but there was no evidence
of additional Tis or papillary tumors anywhere in the
urinary tract. We have commenced an intravesical
treatment course of BCG and MMC.

Side effects
Discomfort, transient frequency, urgency and
occasional chills all occurred and always resolved
within 36 hours. Visible hematuria and fever
exceeding 38°C followed instillations # 5 and # 8
forcing postponement of the subsequent instillations
for 1 week. Six weeks after cessation of therapy the
patient developed urgency, frequency and pelvic pain.
Urinalysis showed marked leucocyturia, slight
hematuria, and routine culture was negative as was
staining for mycobacteria. The prostate specific
antigen (PSA) level increased from 2.2 ng/ml to
7.2 ng/ml. Digital rectal examination and transrectal
ultrasound demonstrated a nodular region in the right
prostatic lobe. Biopsies of the suspicious area showed
granulomatous prostatitis, which slowly resolved
with a 4 month course of isoniazid 300 mg daily and
rifampin 600 mg daily, the PSA level falling to
2.8 ng/ml.

Discussion

It is quite possible that the patient’s initial symptoms
of bladder outlet obstruction were caused by the
irritative effects of multifocal Tis of the bladder.
However, when he presented at our clinic, 2 years later,
hematuria had been present for only 4 months. The
full extent of the carcinomatosis then induced debates
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on different modes of treatment. Diffuse Tis of the bladder alone has considerable invasive potential and the prognosis worsens with associated T1 papillary tumors. Obviously the risks are further increased with concomitant Tis and Ta papillary tumors extending into the prostatic urethra and UUT. The patient’s rejection of extensive surgery, lifelong chronic dialysis and no guarantee of a cure reduced the options to localized immunotherapy or chemotherapy, or both.

BCG is generally accepted as the most effective of the localized instillations for Tis whereas chemotherapeutic agents such as MMC are equally as effective in the treatment of Ta papillary tumors and have less side effects.6 BCG instillations into the UUT have been employed by numerous investigators each with small numbers of patients.7 Some patients have been treated with ureteric instillations of MMC, usually for papillary tumors and initial results were promising.8 Combined treatment with BCG and MMC for active urothelial carcinoma has theoretical appeal. BCG is an immuno-modulator and several induced immunological changes result in destruction of tumor cells. MMC causes synthesis inhibition and strand breakage of DNA. These two distinct mechanisms could be additive and a Finnish group used this combination in patients with bladder Tis, whose results were superior to those treated with MMC alone.9

Most side effects were self limiting and associated with BCG instillations. However, the appearance of a nodule in the prostate associated with a rising PSA level was worrisome, although granulomatous prostatitis following intravesical BCG therapy is well recognized.10

It is difficult to attribute this patient’s extensive disease to his cigarette smoking alone, prolonged and intensive though it was. It is likely that some form of genetic polymorphism made him unduly susceptible to carcinogenic exposure. Unquestionably, apparent elimination of the carcinomatosis with three treatment cycles has not eliminated the cancerous diathesis. This was evident 2 years later with intravesical recurrence and the patient is subject to lifelong surveillance.

References