CASE REPORT

Mycobacterial brain tuberculomas due to Bacille Calmette-Guérin intravesical chemotherapy for bladder cancer: A case report and literature review

Vitaly Golub MD, Prashant Malhotra MD, Shital Patel MD

Bacille Calmette-Guérin (BCG), a live attenuated strain of wild-type Mycobacterium bovis, is widely used for vaccination against tuberculosis. Since 1976, it has been shown to be highly effective in the treatment of superficial bladder cancer (1). Treatment of the latter appears to be relatively safe, with major adverse reactions occurring in fewer than 5% of patients (2). Documented complications of BCG immunotherapy include granulomatous pneumonitis, hepatitis, prostatitis, vascular aneurysms and vertebral osteomyelitis (3).

Central nervous system (CNS) infections caused by M bovis are rare, and those caused by BCG are described in only a few case reports (4-6). CNS infections due to Mycobacterium tuberculosis range from meningitis to tuberculomas, which are tumour-like masses with occasional central necrosis. We describe what we believe is one of the first documented cases of cerebral BCG tuberculoma observed in a 73-year-old immunocompetent man, three years after intravesical BCG immunotherapy. His workup revealed no identifiable extracranial source. He responded well to treatment with rifampin, ethambutol and moxifloxacin.

Patients undergoing intravesical BCG therapy should be closely monitored for the development of this complication. Prolonged antitubercular therapy, possibly including moxifloxacin, appears to be beneficial in the treatment of central nervous system tuberculous infections.

Key Words: BCG; Bladder cancer; Cerebral; Tuberculoma

Des tuberculomes cérébraux mycobactériens causés par une chimiothérapie intravesicale par le bacille de Calmette-Guérin pour le traitement d’un cancer de la vessie : un rapport de cas et une analyse bibliographique

L’immunothérapie par le bacille de Calmette-Guérin (BCG) est largement utilisée pour traiter le cancer superficial de la vessie. Les auteurs pensent que le présent cas est l’un des premiers à rendre compte d’un tuberculome cérébral par le BCG chez un homme immunocompétent de 73 ans, trois ans après une immunothérapie intravesicale par le BCG. Son bilan n’a révélé aucune source extracrânienne repérable. Il a bien réagi au traitement à la rifampicine, à l’éthambutol et à la moxifloxacine. Les patients qui subissent une thérapie intravesicale par le BCG devraient faire l’objet d’un suivi étroit pour décélérer l’apparition de cette complication. Une thérapie antituberculaire prolongée, pouvant inclure de la moxifloxacine, semble bénéfique pour le traitement des infections tuberculeuses du système nerveux central.
remained negative at six weeks. The patient was continued on the above four medications.

Six weeks later, the New York State Department of Health Mycobacteriology laboratory (Wadsworth Center, USA) using an in-house polymerase chain reaction-based genomic deletion analysis (7) for regions of difference (RD) 1, RD9 and RD 10, identified the isolate as an M bovis BCG strain. Sensitivity testing using the Bactec MGIT 960 system (Becton Dickinson, USA) indicated that the strain was resistant to isoniazid (at a concentration of 0.1 µg/mL) and pyrazinamide (at a concentration of 100 µg/mL), but sensitive to rifampin, ethambutol and ciprofloxacin. At this time, isoniazid and pyrazinamide were discontinued and moxifloxacin was started.

One year later, the patient is doing well on ethambutol, moxifloxacin and rifampin. Serial MRIs of the brain have shown a decrease in the size of the cerebral lesions. His right hand tremor has resolved.

**DISCUSSION**

M bovis, the agent responsible for bovine tuberculosis, is also known to cause zoonotic infections. M bovis is a member of the M tuberculosis complex, which includes M tuberculosis, Mycobacterium africanum and Mycobacterium microti. While similar to M tuberculosis, M bovis is uniformly resistant to pyrazinamide, and has a different composition on a nucleic acid probe. Most human exposure stems from raw milk ingestion and inhalation of contaminated droplets (8). In the current era, M bovis constitutes a fraction of human tuberculosis cases. In a 1995 survey from France (8), 0.5% of human tuberculosis cases and, more recently, 1.4% of United States human tuberculosis cases reported (9) were due to M bovis. The vast majority of the patients in the latter study were of Mexican descent. Pulmonary and extrapulmonary manifestations of M bovis are described in the literature. However, the digestive route of acquisition has been largely eliminated with milk pasteurization.

BCG is an attenuated form of wild-type M bovis and has been used for vaccination against human tuberculosis. Since 1976, it has also been used for the treatment of superficial bladder cancer, and approved for the treatment of carcinoma in situ (2). It is believed to exert its antitumour activity via local modulation of immune responses, which results in inflammation and subsequent elimination of malignant cells (10). The most widely used regimen consists of a weekly intravesical BCG injection for six weeks, which is repeated three more times every six months (2).

While generally well tolerated, complications with BCG immunotherapy have been well documented. Most of the early manifestations of BCG disease are characterized by a hypersensitivity type of reaction, namely, pneumonitis, hepatitis or, rarely, sepsis. Fever, chills, sweats and malaise within three months of instillation are the most common presenting symptoms (3). In a Cochrane database review (11) comparing transurethral resection of the tumour with BCG immunotherapy, the most commonly reported complications were fever (25%), cystitis (67%) and hematuria (23%). While the latter reported no deaths due to BCG therapy, fatal sepsis is described in the literature at an approximate rate of one per 15,000 patients (2).

There are no standard guidelines for the treatment of BCG infections occurring as a complication of intravesical therapy. Although most of the early manifestations are believed to be secondary to hypersensitivity, some authors recommend treatment with isoniazid or fluoroquinolone if symptoms persist beyond 48 h (12). In addition, a tapered course of prednisone is occasionally used for hypersensitivity.

Late infections with BCG are characterized by a spectrum of manifestations. In contrast to early disease, which frequently affects more than one organ, late disease is usually localized. Moreover, late manifestations are more likely to yield a positive culture. In a review (11) of BCG infections after bladder cancer treatment, the predominant manifestations were testicular and mycotic aneurysm granulomas. Common to both early and late disease is the pathological diagnosis of a non-necrotizing granuloma. Disseminated or severe infections usually require a prolonged course (ie, six to 12 months) usually with a combination of two or three drugs.

CNS disease caused by M tuberculosis is well described in the literature and includes diffuse exudative leptomeningitis, serous tuberculosis meningitis, epidural or subdural abscess, and intracerebral or intraspinal tuberculosis. Tuberculomas are most frequently encountered in the cerebrum, but those located in the cerebellum are also described and are most common in children. Tuberculomas are usually 2 mm to 12 mm in size, with lobular architecture and commonly central caseous necrosis. The treatment of CNS disease due to M tuberculosis usually involves 12 to 18 months of appropriate therapy.

CNS tuberculosis due to wild-type M bovis have been described in case reports/series (4,13). CNS disease caused by BCG vaccination is extremely rare, with most case reports describing disease only in immunocompromised children (5).

To our knowledge, the present report is the first to describe a case of M bovis BCG causing a CNS tuberculoma in an otherwise immunocompetent patient without any evidence of systemic dissemination after intravesical therapy for bladder cancer. Our patient had received multiple instillations of BCG over a four-year period, the most recent one occurring approximately three years before presentation. Isoniazid resistance was concerning, although we could not ascertain the batch or the strain that the patient received. Isoniazid-resistant BCG lymphadenitis has been rarely described in case reports (14,15).

Our patient responded well to a combination therapy with rifampin, ethambutol and moxifloxacin. A recent study (16) showed that moxifloxacin penetrates well into the CSF. Adequate CSF levels of the fluoroquinolone were recorded in all patients who were involved. Of note, rifampin decreases moxifloxacin concentrations, and dose adjustment may be considered in patients receiving both medications (16,17).

Our case illustrates a rare, but serious infectious complication of intravesical BCG therapy for bladder cancer. Patients undergoing this therapy should be closely monitored for development of these complications. Prolonged antitubercular therapy, possibly including moxifloxacin, appears to be beneficial in the treatment of CNS tuberculous infections.

**REFERENCES**

Submit your manuscripts at http://www.hindawi.com