Voriconazole-Associated Periostitis in a Heart Transplant Patient

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Abstract

A 60 year-old female with a history of a heart transplant was treated with voriconazole for presumed pulmonary aspergillosis. After approximately three months, she began to experience progressively worsening musculoskeletal pain. Her presentation was notable for widespread tenderness, a lack of digital clubbing, and elevated alkaline phosphatase. Bone scintigraphy demonstrated increased activity at multiple locations correlating with the patient’s pain. Periostitis was identified in these locations on radiographs. A few months after discontinuing voriconazole, the patient’s pain had resolved, alkaline phosphatase had normalized, and abnormal activity on scintigraphy had markedly improved.

Introduction

Voriconazole is commonly used for the treatment and prophylaxis of invasive fungal infections in both immunocompetent and immunocompromised patients. Established adverse effects include visual disturbances, hepatic toxicity, and photosensitivity [1]. Recent reports have described painful periostitis in heart and lung transplant patients receiving voriconazole for treatment or prophylaxis of invasive fungal infections.

Case Report

A 60 year-old female with a history of a heart transplant was treated with voriconazole for presumed pulmonary aspergillosis. After approximately three months, she began to experience right arm pain. One month later, she was in a motor vehicle collision, and she attributed subsequent pain to that event. Her pain progressively worsened, spreading to her upper arms, thighs, hips, knees, and hands. Erythrocyte sedimentation rate and C-reactive protein were elevated, but other inflammatory markers including antinuclear antibodies, anti-double-stranded DNA antibody, and anti-histone antibody were negative.

Voriconazole was discontinued after approximately five months, and the patient was hospitalized for pain control and further workup. Rheumatology was consulted for possible voriconazole-associated periostitis. Physical exam demonstrated tenderness to palpation at the shoulders and thighs without synovitis of the hands or digital clubbing. Laboratory results were notable for alkaline phosphatase of 304 units/liter at the time of admission (normal range: 31-95 units/liter). Bone specific alkaline phosphatase was also elevated at 90.8 micrograms/liter (normal range: 5.6-29 micrograms/liter). A technetium-99m Methylene Diphosphonate (MDP) bone scan (Figure 1A), a CT chest (Figure 2A), and a radiographic bone survey (Figure 2B, 2C) were obtained. The bone scan demonstrated multiple foci of cortical MDP activity predominantly at the proximal extremities in a relatively symmetric distribution. Radiographs demonstrated periostitis at these locations, which correlated with the patient’s pain.

When the patient was discharged approximately 2 weeks later, her pain was improving and her alkaline phosphatase had decreased to 217 U/L. Alkaline phosphatase continued to trend down in the subsequent months. Follow-up bone scan (Figure 1B) approximately 5 months after voriconazole was discontinued demonstrated marked improvement in periostitis. By this time, alkaline phosphatase had normalized.

Discussion

Periostitis associated with voriconazole therapy has been reported in patients with lung transplant [2,3], heart transplant [4,5], and other conditions [6]. These patients present with bone pain and elevated alkaline phosphatase. The lack of digital clubbing may help distinguish voriconazole-associated periostitis from secondary hypertrophic osteoarthropathy. Symptoms improve after voriconazole is discontinued, and radiographic and scintigraphic improvement is seen in the subsequent months.

There are two possible theories to explain the development of
periostitis in patients treated with voriconazole. One is based on the fact that fluorine is organically bound in voriconazole and 5% of it is metabolized to free fluoride. It is therefore possible that voriconazole, via hepatic oxidative metabolism, results in unbound fluoride metabolites that cause variably elevated fluoride levels after extended use in predisposed patients via pharmacogenomic variations in drug metabolism. A second possibility is that renal insufficiency predisposes individuals to fluoride accumulation, because its renal clearance is directly related to glomerular filtration rate.

Differently from other reported cases, symptoms can develop shortly after voriconazole initiation. Recognition of this condition is important since discontinuation typically leads to prompt resolution of often disabling pain related to periostitis.

References