Case Report

A Rare and Unusual Case of Burkitt’s Lymphoma Presenting with a Prostate Mass in a 12-Year-Old Boy

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Burkitt’s lymphoma is the most frequent subtype of non-Hodgkin’s lymphoma in childhood. Radiographic findings are protean and can often overlap with other neoplastic and nonneoplastic processes. We present an unusual case of Burkitt’s lymphoma in a 12-year-old boy presenting with a one-week history of urinary retention, dysuria, and “tailbone pain,” as well as a 4-week history of jaw pain, initially treated as a dental abscess. On dental radiography, the patient was found to have resorption of alveolar bone adjacent to the lower first molars bilaterally, in keeping with “floating teeth,” classically associated with Langerhans cell histiocytosis. Additionally, a large, eccentric, prostatic mass was noted, prompting the inclusion of rhabdomyosarcoma on the differential diagnosis, with subsequent definitive diagnosis of Burkitt’s lymphoma on tissue and bone marrow biopsy. This case highlights the imaging overlap of these childhood neoplasms with an unusual lymphomatous prostate mass. It is important that the radiologists and pediatricians be aware of this potential overlap and the unusual presentation of Burkitt’s lymphoma.

1. Introduction

Burkitt’s lymphoma was first described in 1958, by the surgeon Denis Burkitt, who while working in Uganda, noted children with rapidly enlarging tumors of the jaw [1, 2]. The World Health Organization characterizes Burkitt’s lymphoma into 3 types: endemic, sporadic, and immunodeficiency-associated [1]. Endemic Burkitt’s lymphoma is associated with Epstein-Barr virus (EBV) in 95% of cases and is most commonly found in equatorial Africa and Papua New Guinea [1]. The sporadic (or American) type is associated with EBV only 15% of the time, while the immunodeficiency-associated type is seen in patients with HIV, allograft recipients, and those with congenital immunodeficiency [1].

Burkitt’s lymphoma is the most frequent subtype of non-Hodgkin’s lymphoma in childhood, with the jaw and abdomen, specifically terminal ileum, being the most common sites; it grows rapidly, with a doubling time of 24 hours [1]. Jaw involvement is common in the endemic type of Burkitt’s lymphoma but far less common in the sporadic type [3].

2. Case Report

A 12-year-old Caucasian boy presented to the hospital with a four-week history of jaw pain, resulting in difficulty in eating. Upon presentation, he had developed gingivitis and bleeding gums. One week prior to admission, he developed pain in the tailbone area and noted difficulty in urinating with retention symptoms and periodic dysuria. In the week prior to admission, he had been seen at his home hospital emergency room and was started on antibiotics for a presumed dental infection. His family reported an approximate 15-pound weight loss in the month prior to admission. He denied fever but had mild night sweats. He reported low
energy, for which he missed a week of school. He had been otherwise healthy. He takes no medications and reports no allergies and all immunizations were up to date. His past medical history and family history were noncontributory.

Head and neck exam revealed symmetrical swelling around his lower incisors with upwards displacement of the teeth by almost 1.0 cm, both teeth were loose. There was no purulent discharge. There was swelling of the face and lower jaw area, without palpable lymphadenopathy. Note was made of hepatosplenomegaly, as well as bruises on his knees, shin, forearm, and elbow, felt to be from baseball. The physical exam was otherwise normal.

Admission blood work revealed platelets of $33 \times 10^9$ L, WBC $15.9 \times 10^9$ L, and hemoglobin of 141 g/L. Blood smear revealed a left shift with a few circulating blasts and abnormal cells. The liver enzymes were slightly elevated, as was the creatinine at 87 $\mu$mol/L. The lactate dehydrogenase was markedly elevated at $>2500$ U/L, as was the uric acid at 1025 $\mu$mol/L.

Plain film imaging of the jaw performed at an outside institution revealed loss of alveolar bone adjacent to the roots of the lower first molars bilaterally with erosion of the distal roots at these levels, in keeping with “floating teeth” (Figure 1).

An MRI of the brain, face/palate, and pelvis was performed, which revealed multiple lesions within the mandible and maxilla. These lesions were slightly T2 hyperintense and T1 isointense to muscle with homogeneous enhancement. There was involvement of the body of the mandible bilaterally, extending superficially and deeply to the mandibular margins with cortical erosion. Maxillary lesions were also present, extending to the anterior margins of the maxilla, again with cortical erosion (Figures 2 and 3). The brain and pituitary gland were normal.

Within the pelvis, there was a well-circumscribed, periurethral mass within the left lobe of the prostate gland ($3 \times 4 \times 4$ cm) demonstrating slight T2 hyperintensity to muscle with irregular, linear central hypointense regions, and faint enhancement on the postcontrast T1 FS images. As well, a 2 cm mass at the superior/posterior aspect of the bladder on the right (Figures 4, 5(a), and 5(b)) was identified, demonstrating similar signal characteristics. Hepatosplenomegaly was also noted. A contrast enhanced CT of the chest, abdomen, and pelvis demonstrated multiple, bilateral subcentimeter hypodense lesions involving both kidneys (Figure 6). On CT, the prostate demonstrated asymmetric...
A right mandibular biopsy showed diffuse infiltrate of medium to large lymphoid cells that were monomorphic against a heavy background of scattered macrophages with a “starry sky” pattern (Figure 7). Bone marrow biopsy from the right iliac crest showed more than 95% pattern of infiltration with Burkitt lymphoma, which was confirmed with immunophenotyping of the neoplastic cells in the marrow. Cerebrospinal fluid showed clusters of cells with degeneration that were similar to the known neoplastic cells.

The patient was diagnosed with stage IV Burkitt’s lymphoma and COP reduction chemotherapy was initiated immediately. The patient experienced a dramatic reduction in his tumor burden, with follow-up imaging of the prostate revealing near complete resolution of the mass. Unfortunately, the patient relapsed with Burkitt’s Leukemia approximately 6 months after initial treatment was started, with 99% blast involvement of his bone marrow. During ongoing therapy, the patient experienced sepsis in the context of profound pancytopenia, acute kidney injury requiring continuous renal replacement therapy, enterococcus pneumonia requiring intubation, and a large pericardial effusion. On day 15 after cycle number 2 of his chemotherapeutic regime for Burkitt’s leukemia, he developed severe lactic acidosis, respiratory failure, and severe, profound bradycardia that could not be reversed. He died the following morning, seven and a half months from his initial diagnosis.

3. Discussion

Though the most common type of non-Hodgkin’s lymphoma in children, prostate involvement of Burkitt’s lymphoma is uncommon and accounts for <0.1% of genitourinary involvement [4]. In a multi-institutional study of 62 cases of malignant lymphoma involving the prostate, only one case was found to be Burkitt’s lymphoma; this happened to be in the single child [5]. In this case series, there was a 5-year-old boy with secondary involvement of the prostate by Burkitt’s lymphoma, who died 1 week after diagnosis. The imaging findings in this case were not described.
The terminal ileum is the most common location of Burkitt’s lymphoma in children [1]. Though abdominal and pelvic involvement are common, prostatic involvement of Burkitt’s lymphoma, specifically in children, has not been previously described in the imaging literature. In their report of 62 cases of malignant lymphoma involving the prostate, predominantly in adults, Bostwick et al. found that secondary involvement of the prostate was more common than primary involvement (65% versus 35%), and that lymphoma specific survival was 64% at 1 year and 50% at 2 years [5].

Specific imaging of the prostate gland is rarely warranted in children but is included during workup of symptoms related to the lower genitourinary tract, including urinary retention, hematuria, dysuria, and incontinence, or during investigations for suspected congenital anomalies [6]. In children presenting for workup of a prostatic mass, rhabdomyosarcoma would be high on the differential diagnosis, as it is the most common tumor of the lower genitourinary tract in children and often involves the prostate gland [6, 7]. Other pediatric prostatic tumors are extremely rare [5, 8, 9]. These children often present with symptoms of urinary and fecal retention. Rhabdomyosarcomas originating in the prostate carry significantly worse prognosis than do tumors that involve bladder only [8]. Bladder wall invasion may be detected on MR imaging, with T2-weighted images demonstrating higher signal intensity tumor extending into the lower-signal intensity wall. Perivesical and perirectal fat invasion can be demonstrated on T1-weighted images [6].

Leukemic infiltration has a similar MR appearance to lymphoma, with hypovascularity and only mild contrast enhancement [6]. A case of myeloid sarcoma of the prostate in a child with acute myelogenous leukemia has been reported [9]. MR imaging was not performed in this case, with sonography showing a hypochoic mass involving the left bladder wall displacing the rectum posteriorly. This lesion was irregular and heterogeneous on contrast enhanced CT [9].

Prostatic carcinoma and carcinoid have also been reported in the pediatric population [10, 11]. A case of primary carcinoid of the prostate in a 7-year-old boy with multiple endocrine neoplasia IIb has been reported, described as a T2 hyperintense mass without extension beyond the prostate [11]. Inflammatory tumor of the prostate in a child has been reported as a cystic lobular and centrally necrotic midline tumor that nearly completely resolved with antibiotic therapy [12]. Chronic prostatitis is rare and may be secondary to abnormal voiding conditions [6].

As previously mentioned, jaw involvement with Burkitt’s lymphoma is more commonly seen with the endemic form though a case describing “floating teeth” at presentation in sporadic Burkitt’s lymphoma in a 66-year-old male has been published [13]. Historically, “floating teeth” in pediatric patients have been thought to be almost pathognomonic of Langerhans cell histiocytosis, though this finding may also reflect any destructive process in the mandible, including infectious, hematologic, metabolic, or neoplastic etiologies [14]. In our case, the presence of floating teeth on plain radiography prompted not only the MR imaging of the jaw but also the pituitary, which can also be involved in Langerhans cell histiocytosis and is often associated with diabetes insipidus [15].

Though a definitive diagnosis was swiftly made on tissue and bone marrow biopsy, this case highlights the imaging overlap of these childhood neoplasms. It is important that the radiologist and pediatrician be aware of these similarities and that not all pediatric prostatic masses reflect rhabdomyosarcoma. Definitive diagnosis requires histologic examination in all cases.

**Conflict of Interests**

The authors declare that there is no conflict of interests regarding the publication of this paper.

**References**


