An Interesting Case Presentation of Burkitt Lymphoma Presenting with Lactic Acidosis and Thrombocytopenia

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Received: 03 Apr, 2020 | Accepted: 27 Apr, 2020 | Published: 01 May, 2020

Citation: Miah A (2020) An Interesting Case Presentation of Burkitt Lymphoma Presenting with Lactic Acidosis and Thrombocytopenia. J Clin Case Stu 5(2): dx.doi.org/10.16966/2471-4925.201

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Abstract

Burkitt Lymphoma (BL) is a very aggressive B-cell lymphoma caused by the translocation and subsequent dysregulation of the c-Myc gene on chromosome 8. There are 3 distinct clinical forms of Burkitt Lymphoma: endemic (African), sporadic (non-endemic), and immunodeficiency associated. Although these forms differ in their clinical presentation and their epidemiology, they share the same aggressive clinic behavior and are histological identical. This case presentation reports a rare case of sporadic form of BL clinically manifesting as sepsis in an immunocompetent young adult male. A quick diagnosis with bone marrow biopsy leads to starting treatment with IV chemotherapy. Unfortunately, our patient succumbed to the disease. It is necessary to consider malignancies as one of the main differential diagnosis to facilitate the correct diagnosis and prompt treatment.

Keywords: Burkitt lymphoma; Non-Hodgkin lymphoma; Hyper-CVAD; Sepsis; Lactic acidosis

Abbreviations: BL: Burkitt Lymphoma; IVDU: Intravenous Drug Use; DIC: Disseminated Intravascular Coagulation; TTE: Trans Thoracic Echocardiogram; ID: Infectious Disease; VBG: Venereal Blood Gas; CRP: C-Reactive Protein; UDS: Urine Drug Screen; UA: Urine Analysis; BUN: Blood Urea Nitrogen; VDRL: Venereal Disease Research Laboratory; PPD: Purified Protein Derivative; LDH: Lactate Dehydrogenase; PT: Prothrombin Time; INR: International Normalized Ratio

Introduction

Burkitt Lymphoma (BL) is a rare and highly aggressive B cell neoplasm comprises 30 percent of non-endemic pediatric lymphomas, but less than 1 percent of adult non-Hodgkin lymphomas [1,2]. It is endemic in Africa and sporadic in other parts of the world and is usually diagnosed in children and young adults but rarely in middle-aged adults [3]. Patients with BL present with rapidly growing tumor masses and often have evidence of tumor lysis, however sometimes they can present in unusual ways. We present a case of a 29-year old male who presented with tachycardia, persistent lactic acidosis and DIC. The acidosis was so severe and persistent that the patient required intubation from respiratory fatigue and even required dialysis. In addition, despite being on broad spectrum antibiotics he continued to have fevers.

After his workup did not support any of the more common causes of these symptoms, such as negative cultures, further investigation with bone marrow biopsy revealed Burkitt lymphoma. He was started on aggressive treatment with hyper-CVAD alternating with high dose Methotrexate and high dose Cytarabine. Unfortunately, his malignancy was so aggressive and included CNS involvement which leads to tracheostomy placement, paralysis, and eventually death. This is an example of a rare condition with an even more rare presentation. This diagnosis was not reached until many more common diagnoses were ruled out. It is an example of a case where a broad differential is needed to secure the correct diagnosis.

BL is caused by the translocation and subsequent dysregulation of the c-Myc gene on chromosome 8 [4]. There are 3 distinct clinical forms of Burkitt lymphoma: Endemic (African), Sporadic (non-endemic), and Immunodeficiency Associated [5]. Although these forms differ in their clinical presentation and their epidemiology, they share the same aggressive clinic behavior and are histological identical [6]. The sporadic form is the most common in USA and is identified in this case and represents 30% of pediatric non-endemic lymphomas and less than 1% of adult non-Hodgkin lymphomas [7]. However; the clinical presentation of Burkitt's lymphoma will differ depending upon the specific variant. In endemic areas, it usually involves the facial bones, particularly the jaw, maxilla, and orbit, especially in young children. This variant has been associated with Epstein-Barr Virus (EBV) infection, as well as frequent concomitant malaria infection [8]. In comparison, the sporadic form tends to present in the lymphoid tissues of the gut, often presenting as masses

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in the terminal ileum, or even with massive abdominal involvement. Bone marrow involvement is commonly seen in progressive disease [9]. EBV involvement is reported in around 15-30% of cases [10]. The immunodeficient form is often associated with HIV infection, and may also be seen in post-transplant patients who are chronically immunosuppressed [9].

Case Presentation

A 29-year old white male presented with bilateral hip and back pain. His past medical history was significant for IVDU. Initially, the symptoms started several weeks ago prior to admission when he began having left hip pain which progressively became worse. He stated the pain spread to involve both hips and he rated it (10/10) and described it as sharp. Any type of movement makes the pain worse. He states that the pain progressed to involve his lower back and described it as throbbing. He also stated that he is having some left lower face tingling. Otherwise he denies any numbness and tingling in lower extremities. He tried taking over the counter Aleve which did not alleviate his pain. Aleve and Ibuprofen were also ineffective at treating the pain. He denies any sick contacts, recent IVDU, including injection into his hip. He did not take any Tylenol to treat the pain. He denies saddle paresthesia or bowel or bladder incontinence. He reports no focal infiltrates and an HIV antibody screen was negative, as was B, C, and HIV serologies negative, Albumin 3.2; Alkaline phosphate 255; AST-267; ALT-112; Total bilirubin-1.3, Direct bilirubin-0.7, PT-34, Na-133; K-4.1; Cl-96, CO$_2$-15, AG-22; Ca-9.7; Glucose-45; BUN-16, Cr-0.69, CRP-253, ESR-26, UA: Ketones; UDS: Suboxone and Oxycodone, LDH was too high to be detected, Hepatitis B, C, and HIV serologies negative, Albumin 3.2; Alkaline phosphate 255; AST-267; ALT-112; Total bilirubin-1.3, Direct bilirubin-0.7, PT-16.7; INR-1.6; Fibrinogen-311. A chest X-ray, PA and lateral showed no focal infiltrates and an HIV antibody screen was negative, as was a later PCR. CT Chest abdomen and pelvis showed splenomegaly, otherwise within normal limits. Patient was admitted with sepsis suspicion for endocarditis.

After admission, the patient spiked episodic fevers as high as 38.8°C. He was transferred to ICU service a day later. Since transfer, his lactate had increased from 11.1 to 13.2 and he became tachypneic. ID was consulted and recommended starting him on broad spectrum antibiotics, he was also started on a sodium bicarbonate drip, and he eventually needed intubation for respiratory compromise. He needed emergent dialysis for overwhelming acidosis. TTE with pericardial effusion, no vegetations. MRI brain was within normal limits. Hematology/Oncology was consulted for thrombocytopenia. Initially they were concerned about DIC given his history of IVDU.

Blood and urine cultures were drawn on admission. The following day, the patient continued to have episodic fevers and he also became hypertensive with a BP 190/90. The blood cultures drawn on admission showed no growth in four days. Further workup including ADAMTS13 did not reveal cause of persistent thrombocytopenia. A peripheral smear was performed which showed schistocytes, normochromic normocytic anemia, severe thrombocytopenia, banded neutrophils, myelocytes, RBC inclusion bodies (Figure 1).

Bone marrow was biopsy performed 4 days after admission; the results were consistent with Burkitt Lymphoma findings with the pathognomonic “starry sky” appearance (Figure 2). It showed a hypercellular marrow (greater than 95% cellularity), flow cytometry was CD10-positive B-Cell lymphoma (76%), and karyotype: 8,22 translocation.

Throughout his hospital course, patient continued to experience episodic fevers as well as uncontrolled hypertension. Patient also became confused and would regularly only be oriented to person and/ or place. An ophthalmologic exam revealed papilledema. Multiple escalations of his blood pressure regimen failed to bring his pressures within reasonable limits. Patient underwent a traumatic lumbar

**Figure 1:** Peripheral smear showed schistocytes, normochromic normocytic anemia, severe thrombocytopenia, banded neutrophils, myelocytes, RBC inclusion bodies.
The staging workup must be expedited, because Burkitt tumor grows rapidly, causing life-threatening complications. BL is classified as a highly aggressive peripheral B cell tumor and demonstrating the highest proliferation rate of any neoplasms in humans, with a potential doubling time of 24 hour and a growth fraction of nearly 100% [11]. Sudden and unexpected development of life threatening complications such as airway and abdominal obstruction and acute renal failure has been reported with Burkitt lymphoma in the literature [12]. The diagnosis of Burkitt lymphoma is made by obtaining a biopsy of the tumor mass for histopathology, immunohistochemistry, and flow cytometry cytogenetic studies to identify c-Myc mutation will aid in the diagnosis. The most suggestive lymph nodes should be selected for excisional biopsy. c-Myc translocation is a characteristic of Burkitt lymphomas. Approximately 80% of Burkitt lymphoma cases have the t(8;12) translocations. An increase in c-Myc expression results from the translocation. This c-Myc dysregulation seems to be the defining abnormality that eventuates into BL [13,14]. They are recognized as small non-cleaved cell lymphomas displaying a starry sky appearance due to the high rate of proliferation and spontaneous cell death (Figure 2).

This case presents a very rare type of presentation of a very rare type of lymphoma in adults. Usually when a young patient like in our present study with fever we think of sepsis. This is an important case because his initial presentation was similar to sepsis and his history of IVDU led us towards a diagnosis of endocarditis. Sepsis will always remain high on the differential diagnosis, however if they do not respond appropriate antibiotics and if cultures are negative other diagnosis need to be considered. One unique presentation that our patient had was severe lactic acidosis and LDH which was not mentioned in other case presentation. A possible explanation could be due to high tumor burden seen in BL similar to a tumor lysis syndrome presentation. A broad differential with alternative diagnoses must be considered in patients who fail to improve with appropriate therapy. In literature review, most cases of BL do not present with lactic acidosis and it is considered a rare complication of BL with only 26 previous reports [15]. However, with those cases that present with lactic acidosis, the mortality rate is at 73% at 1 month [15]. A possible explanation is that with aggressive lymphoma lactic acidosis is common due to overproduction of lactic acid by rapidly dividing tumor similar to what is seen in BL.

References

Discussion and Conclusion
Burkitt lymphoma is an extremely rapidly growing tumor with a high sensitivity to chemotherapy but in which drug resistance can develop quickly [9]. Although Burkitt lymphoma is one of the most aggressive types of cancer it is also one of the most curable [7]. Chemotherapy is the mainstay of treatment for Burkitt lymphoma. Treatment of limited stage Burkitt lymphomas is usually very successful in children, with a cure rate of over 90%. For more advanced Burkitt lymphoma cure rate ranges from about 80% to 90%. However, compared to adults with low risk the five-year survival is 70% and high risk cases the survival is less than 30% [4].

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References


