Synchronous Burkitt’s Lymphoma of Breast and Colon in a Female Patient: A Report of a Case and Review of Literature

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SUMMARY
Burkitt’s lymphoma (BL) is a highly aggressive cancer of B-cells, which is more prevalent in African children. Extranodal involvement of the BL is not a common entity. Herein, we report a young woman who referred to University hospital due to rectorrhagia and breast swelling. Colonoscopy examination revealed a large tumoral lesion of 5×6 cm with superficial ulceration in proximal of ascending colon. Computed tomography (CT) scan of the abdomen showed a 55×42 mm mass in ascending colon and caecum. No lymphadenopathy, splenomegaly and hepatomegaly were reported. Breast ultrasound was performed showed a 24×11 mm hypoechoic heterogeneous mass in the left breast. Bone marrow biopsy was normal. Immunohistochemical assay of colon and breast biopsies confirmed the diagnosis of primary BL. After four courses of chemotherapy, PET-CT revealed complete remission and no sign of metastasis. Since BL is a rapidly-growing neoplasm, prompt diagnosis and treatment can prevent unnecessary surgery and consequences of delay in treatment.

Keywords: breast neoplasm; Burkitt lymphoma; colon neoplasm; PET-CT.

Introduction
Burkitt’s lymphoma (BL) is a B-cell malignancy in non-Hodgkin lymphoma category. BL is a rare but aggressive cancer that mainly affects African young children associated with Epstein–Barr virus infection as endemic forms. It also may present concerning immunodeficiency in HIV positive subjects or organ transplant recipients.[1] Sporadic variant of the BL is not specific to a certain geographical region and often involves gastrointestinal and respiratory system.[2] Lymphoma represents a rare type of primary breast malignancy (less than 1%) with diffuse large B-cell lymphoma (DLBCL) as the most common; BL of the breast is extremely unusual.[3] We report a rare case of BL with ascending colon and breast involvement presenting with rectorrhagia and unilateral breast mass.

Case Report
A 29 years-old woman presented with a 40-days history of abdominal pain and diarrhea to University clinic. At attendance, there was a complaint of rectorrhagia.
The other symptoms were nausea, changes in bowel habits and weight loss, the sensation of breast mass since two weeks before her admission to the hospital. There was no history suggestive of any fever and night sweats. She had a 3-year-old child and was not pregnant or breast-feeding at the time of referral.

Her routine investigations revealed hemoglobin of 11.7, white cell count of 3100/mm³, platelet count of 240000/mm³, erythrocyte sedimentation rate (ESR) was 19 mm, uric acid was 10 mg/dl, liver functional tests and electrolytes were within normal range and beta2 microglobulin was 1.47.

Colonoscopy examination revealed a large tumoral lesion of 5×6 cm with superficial ulceration in proximal of ascending colon. Computed tomography (CT) scan of the abdomen showed a 55×42 mm mass in ascending colon and caecum. No lymphadenopathy, no splenomegaly and no hepatomegaly. Endoscopic ultrasonography was normal. Breast ultrasound was performed, which showed a 24×11 mm hypoechoic heterogeneous mass in the left breast. Color Doppler revealed peripheral and central vascularity. Histopathological examination of colonic and breast biopsies revealed a malignant neoplasm with ulceration composed of fibrin deposition, cell debris and neutrophilic infiltration. Extensive infiltration of atypical lymphocytes with hyperchromatic nuclei, regular borders and scattered cytoplasm were observed. Also, a few mitotic figures and a segment of necrotic tissue were noted. No granular structures and epithelial cells were reported.

Bone marrow biopsy was normal and immunohistochemistry (IHC) result was as follows: CD79a positive in background B-cells, PAX5 and CD38 positive in neoplastic B-cells, CD43, CD5 and CD3 positive in background T-cells, CD45 positive diffusely, BCL6 positive, CD23, cyclin D1, CD56, MNF116, TdT, CD99 and chromogranin were negative in tumor cells. MYC was positive in more than 80% of the tumor cells and ki67 was positive in more than 95% of tumor cells, so the findings were consistent with Burkitt’s lymphoma.

Soon, she received immune-chemotherapy using Rituximab and Hyper-CVAD (Cyclophosphamide, Vin-cristine, Adriamycin, and Dexamethasone) regimen. Since the closest time to schedule a PET scan was one month later (due to limitations) and because of disease nature, it was not possible to order it before chemotherapy. One month later, PET scan revealed no evidence in favor of metabolically active lesion throughout the suggesting a complete remission of the disease (Fig. 1). Patient’s first visit was in August 2019 and the last follow-up was in January 2020, when the patient showed no sign of disease progression. The patient's consent was obtained for this case study.

Discussion

In 1958, Irish surgeon Denis Burkitt was first to describe BL as an unusual disease in African children.[4] While the exact etiology of BL remains unknown, several risk factors have been introduced. EBV is a human herpes virus (HHV4) with a worldwide spread that most adults are seropositive for it. This B-lymphotropic virus induces B-cell proliferation and may establish latent infection. Transformation of B-cells is mediated by viral antigens, including latent membrane proteins, nuclear antigens, non-coding and micro-RNAs. Hodgkin lymphoma (HL), diffuse large B-cell lymphoma and BL, are among B-cell neoplasms to be associated with EBV infection.[5,6] Alteration and translocation of c-myc proto-oncogene (located on chromosome 8) is a hallmark presentation of BL. Myc acts as a transcription factor to control many cellular functions, including cell growth and proliferation, protein synthesis and apoptosis.[7] Overexpression of the Myc using Immunoglobulin heavy chain enhancer [t(8;14)] would lead to lymphoid system malignancies, including BL.[8] In endemic BL, there is also evidence of the role of Plasmodium falciparum infection in DNA damage modulated by activation-induced cytidine deaminase (AID) in this type of malignancy.[9] BL is a rapidly growing high-grade cancer. Sporadic type BL is more common in western regions and children with male predominance.[10] Up to 40% of extranodal involvements of lymphomas occur in the gastrointestinal system.[11] According to a study concerning children with gastrointestinal BL,
symptoms according to frequency were reported as follows: abdominal pain and swelling, vomiting, intestinal obstruction, constipation, diarrhea, melena and rectal bleeding.[1] Therefore, lower GI bleeding in our patient may be considered as a rare symptom of BL.[12] Intussusception and obstruction of bowel can also mimic acute abdomen, which requires prompt surgical intervention. Small intestine and ileoceleal region are common sites to be affected.[13]

Only approximately 0.5% of breast cancers are attributed to primary lymphoma. This rare manifestation may occur in uni- and bilateral forms. DLBCL accounts as the first prevalent subtype of primary lymphomas of the breast followed by follicular lymphoma, mucosa-associated lymphoid tissue lymphoma and BL.[14] Usually, cases are pregnant or in lactation period; therefore, the diagnosis may be missed due to suspicion of abscess.[14,15] Also, breast BL is reported in association with HIV infection too.[16] Breast swelling, progressively increasing mass, pain, breast enlargement and tenderness are among man manifestation which is not different from carcinoma symptoms.[14,16-18]

The 5-year survival of BL from 43% in 1973-2001 reached 56% in 2002-2008; however, older patients tend to have poorer survival.[19] It has also been shown that risk factors for low survival differ in children and in adults. In adults primary tumors in bone marrow, stage III and IV and in children stage IV were associated with worse survival.[20]

Also, breast BL lacks pathognomonic features in imaging modalities, including mammography and ultrasound.[21]

There is still no treatment of choice for the BL. The effectiveness of surgery is a controversial and usually dramatic response to chemotherapy is observed with complete remission.[22] As BL is a rare but aggressive entity, prompt diagnosis and rapid initiation of treatment can play a substantial role in the control and remission of it.

Informed consent: Informed consent was obtained from the patient in this study.

Peer-review: Externally peer-reviewed.

Conflict of Interest: No conflict of interest.


References


