Generalised Burkitt’s lymphoma involving both breasts - A case report

*S. O. Fadiira, **V. O. Mabayoje, *A. O. A. Ademowo, *M. L. Adeoti,  
**S. A. Olatoke and *A. S. Oguntola  
*Department of Surgery, **Department of Haematology  
LAUTECH College of Health Sciences, P. M. B. 4400, Osogbo.  
E-mail: sofadiira@yahoo.com

Summary
Background: Burkitt’s lymphoma is a disease of children age ranging 8-10 years. Lymphoma involving the breast is an unusual clinical entity, which is rarely distinguished preoperatively from other more common forms of breast cancer. The management differs from the more typical adenocarcinoma of the breast in that the emphasis is on systemic therapy.

Case: We report a 27-year-old pregnant Nigerian civil servant at 28 weeks gestation, who presented with multiple organ swellings including both breasts. She was diagnosed histologically as primary breast Burkitt’s lymphoma. She was treated with systemic chemotherapy after spontaneous abortion. She had a recurrence of the lesion in the right breast a month later that was excised. Presently there is no evidence of any mass.

Conclusion: Compared with breast carcinoma, primary breast lymphoma is a rare disease but should be considered in the differential diagnosis of breast masses.

Key-words: Generalised Burkitt’s lymphoma, Breast, Pregnancy.

Résumé
Introduction: Lymphome de Burkitt est une maladie chez des enfants avec la tranche d’âge de 8-10 ans. Lymphome impliquant le sein est une entité clinique peu commen, ce qui est rarement reconnaisable prééventuellement par rapport au plus adnocaenome typique du sein dans la mesure où l’accent est mis sur la thérapie systémique.

Cas: Nous rapportons le cas d’une fonctionnaire nigériane enceinte âgée de 27 ans pendant une période de 28ème semaine de gestation qui s’est présentée atteinte de la tuméfaction multiple d’organe y compris les deux seins. On l’avait été diagnostiqué histologiquement comme lymphome de Burkitt de sein primaire. On l’avait traité avec la chimiothérapie systémique après une avortement volontaire. Elle avait eu une recuentesse de la lésion dans le sein du côde droit un mois après une intervention chirurgicale. A présent, il n’y a aucun signe de n’importe quel gestelement.

Conclusion: Par rapport au carcinome du sein, lymphome du sein primaire est une maladie rare mais devrait être considéré dans le diagnostic différentiel des masses du sein.

Introduction
Lymphoma involving the breast, either as a primary site or as a site of recurrence from lymphoma previously diagnosed elsewhere is rare. Primary breast lymphoma is seldom distinguished preoperatively from other more common forms of breast cancer. It constitutes 0.04% to 0.5% of malignant breast neoplasms; 1.7% of all extranodal Non-Hodgkins lymphomas, and 0.7% of all Non-hodgkin Lymphomas; in one of the several series.

Breast lymphoma usually manifests as painless masses incidentally detected by self examination or during routine mammography. The majority of the cases are unilateral and of B-lymphocyte histologic lineage; while bilateral involvement is seen almost exclusively in secondary cases and in Burkitt’s lymphoma.13 Prognosis depends on the histological type and stage at diagnosis.45 Multiorgan involvement is commonly seen and secondary cases with mucosal associated lymphoid-tissue (MALT) type lymphoma appear to have a better prognosis.13,8

The purpose of this paper is to consider Burkitt’s lymphoma as a differential diagnosis of breast mass though very rare and to highlight its peculiar presentation.

Case report
A 27-year-old Nigerian Civil Servant presented with 2 months history of painless multiple body swellings involving both breasts, lower abdomen and the back, with associated weight loss, early satiety and vomiting of recently eaten food.

Examination revealed a young lady pale, with evidence of weight loss. There was a left breast mass of 44 cm by 32 cm while the right side was 46 cm by 30 cm. (see fig. 1)

Abdominal examination revealed right iliac fossa mass of 8 cm by 5 cm size, and left lumbar mass of 15 cm by 17 cm, fundal height was at 28 weeks gestation. The last menstrual period (L.M.P) was 12/11/2002; the expected date of delivery (EDD) was 19/8/2003; with viable fetus. An abdominal ultrasound (USG) confirmed the pregnancy and the gestational age in addition to the multiple abdominal masses, a diagnosis of malignant lymphoma in pregnancy was made.

The patient had incision biopsy on the left breast mass, and an aspiration biopsy on both breasts, incisional biopsy of the right breast confirmed the diagnosis (see fig.2). She had a bone marrow aspiration which was negative for...
Burkitt’s cells.

Laboratory values were Packed Cell volume (PCV) 22%; total white blood cell count of 8.7 x 10^9/L; a differential count of Neutrophils 86%, eosinophils 2% and lymphocytes 12%. The ESR was 81mm in the first hour and the alkaline phosphatase was raised at 210 U/L. Other biochemical findings were essentially normal but for elevated urea of 7.6 mmol/L.

A definitive diagnosis of Burkitt’s Lymphoma (stage D due to breast involvement) in pregnancy was made she was scheduled to commence chemotherapy with I.V. Vincristine, she however had a spontaneous abortion before the commencement hence, she was placed on full regimen for Burkitt’s lymphoma as follows: -

- I.V. Cyclophosphamide 1gm/m^2 day 1
- I.V. Methotrexate 37.5mg/m^2 day 1
- I.V. Vincristine 1.5mg/m^2 day 1

The intravenous drugs were given on day one of a fourteen-day cycle.

Oral prednisolone 20mg 8hrly for 5 days with Mesi.
Magnesium trisilicate 15mls 8 hrly also for five days; Allopurinol 100mg orally 8 hrly daily for Fourteen days within a cycle was also given.

She received seven cycles of chemotherapy including (intrathecal central nervous system (CNS) prophylaxis to prevent CNS involvement) with complete regression of all the swellings; A month later she presented with right breast recurrence that was excised. Presently she has no breast mass.

She is currently attending both surgical and haematology clinics for routine follow up. Haemogram pattern of the last follow up visit revealed packed cell volume (PCV) of 29%; total white cell count of 7.6 x 10^9/L; platelet count of 120 x 10^9/L with a white cell differential of Neutrophils 46%. Eosinophils of 14% lymphocytes of 40%, and ESR of 15mm in the first hour. She is presently being screened for possible parasitic infestation though bone marrow eosinophilia is not an unusual finding in malignant lymphoma. The patient tested negative for HIV/AIDS.

Discussion

Burkitt’s lymphoma in this part of the world is usually seen in children, age ranging between 8-10 years, however the case being reported is a 27-year-old pregnant woman. Well known predisposing factors to the development of the disease include residing in malaria endemic areas, and the rain fall pattern with high humidity associated with equatorial forest zones all of which are present in the south west area of Nigeria where the patient resides. Infection with Epstein Barr virus is also a known risk factor, but it is as of now not possible for us to prove that this patient is harbouring the virus.

Non-Hodgkins lymphoma of the breast is uncommon with variable incidence of primary and secondary cases. The largest individual series with clinical follow up has been reported by Giardini et al. In this series thirty-five patients presenting with stage I or II primary breast lymphoma were identified over a 30-year period, with the most common histology as diffuse large cell lymphoma (17 patients) and only 7 patients having low grade malignancy.

A variety of single and combined modality of treatment were used, with a mean follow up of 45 months, seventeen patients had died of their disease, an overall 5-year survival of 43%.

A preponderance of right-sided lesions has been reported while bilateral involvement, indicates secondary involvement. Diffuse large cell lymphoma of B-lymphocyte lineage has been the most frequent histologic subtype in both primary and secondary cases.

Our patient has multiple painless organ involvement including the breast which was painless at the initial presentation, however in one case of primary T-cell lymphoma of the breast reported by Cohen and Brooks, the patient had breast pain and skin breakdown. Breast swelling as a presenting feature of lymphoma has been reported, but breast pain was not mentioned as a predominant feature in these cases. Inalsingh reported a case initially diagnosed as mastitis, but eventually the patient had a biopsy to rule out malignancy, and a histopathologic diagnosis of primary lymphoma of the breast was made. Rarely bilateral inflammatory Carcinoma of the breast may present with fluid features of inflammation; Burkitt’s lymphoma involving the breast, especially in lactating women can also manifest as a rapidly enlarging mass with features of inflammation.
Mammography is indicated in clinically suggestive cases, however, mammography is not reliable for diagnosing breast lymphoma. Needle aspiration or excisional biopsy is necessary to establish a diagnosis. It is necessary to take samples from both breasts because primary breast lymphoma and contralateral breast cancer had been reported. Treatment options include combination chemotherapy such as cyclophosphamide, doxorubicin, vincristine, and prednisone, radiation, and surgical resection.

References