Lupus: The Great Pretender, Breast Manifestations of SLE

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Abstract

Lupus mastitis is an infrequent sequela of the autoimmune disease Systemic Lupus Erythematosus (SLE) which can be confounding to both clinicians and diagnosticians due to its similar appearance to an infectious or even neoplastic process. This case demonstrates a patient who presented with a painful and edematous breast that was initially concerning for mastitis or underlying abscess on the initial clinical examination. After a trial of antibiotics, the patient’s symptoms minimally improved with subsequent work up using mammogram and ultrasound concerning for a possible malignancy. Ultrasound guided needle and core biopsies eventually led to the diagnosis of lupus mastitis. This report discusses the clinical, radiological and histological presentation of lupus mastitis. This is importation for clinicians so they can have a better understanding of this unique presentation of SLE and can prevent unnecessary invasive procedures which may further increase inflammation involved with lupus mastitis. While SLE has a multitude of clinical presentations this case report demonstrates another unique way that SLE can be encountered.

Introduction

In October 2014, a 39 year old female with PMH of Crohn’s disease, Systemic Lupus Erythematosus (SLE), Chronic Heart Failure (CHF) and Hypertension (HTN) was admitted to our hospital due to a progressively painful and edematous right breast. The symptoms lasted for approximately 2 weeks and they were concerning for mastitis or underlying abscess on the initial clinical examination. After a trial of antibiotics, the patient’s symptoms minimally improved with subsequent work up using mammogram and ultrasound concerning for a possible malignancy. Ultrasound guided needle and core biopsies eventually led to the diagnosis of lupus mastitis. This report discusses the clinical, radiological and histological presentation of lupus mastitis. This is importation for clinicians so they can have a better understanding of this unique presentation of SLE and can prevent unnecessary invasive procedures which may further increase inflammation involved with lupus mastitis. While SLE has a multitude of clinical presentations this case report demonstrates another unique way that SLE can be encountered.

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Figure 1: (a). RMLO (b). LMLO (c). RCC (d). LCC. Bilateral MLO and CC views show increased skin and trabecular thickening throughout the right breast as compared to the left.

Targeted right breast ultrasound demonstrated diffuse edematous breast tissue (Fig 2a) with a questionable solid component (Fig 2b, red arrow) and right axillary lymphadenopathy (Fig 2c).

Given that these findings persisted with antibiotic treatment, an underlying malignancy was included in the differential diagnosis, and the examination was given a BIRADS 4 rating which warranted a biopsy.

Ultrasound guided biopsies were performed targeting the solid component noted in the edematous breast tissue (Fig 3a) and right axillary lymphadenopathy (Fig 3b).

Figure 2: (a/b). Edematous retroareolar tissue in the right breast with heterogeneous echotexture, and a questionable solid component. (c). Right axilla demonstrating an enlarged and abnormal appearing lymph node.

The pathology results from the ultrasound guided biopsy of the right breast were reported as: benign breast parenchyma with fibrocystic changes including ductal dilatation, stromal fibrosis, sclerosing adenosis and mild perilobular and intralobular lymphoplasmacytic infiltrate (Fig. 4a). No acute inflammatory changes or malignant cells were noted. The right axillary lymph node showed reactive benign lymphadenopathy. These pathology results were considered discordant since the radiographic findings were still concerning for an underlying inflammatory breast cancer. Consequently, a skin punch biopsy was recommended for a definitive diagnosis.

The skin punch biopsy (Fig 4) demonstrated superficial and deep perivascular lymphoplasmocytic infiltrate (Fig 4b), consistent with a history of systemic lupus involvement with perivascular and periadnexal lymphoplasmacytic infiltrate (Figure 4c/4d); no malignant cells were visualized.
Lupus mastitis typically presents as a tender mass on clinical exam. This disease is not a neoplastic process, but rather, an infrequent sequela of the autoimmune disease systemic lupus erythematosus (SLE). SLE has the potential to involve a variety of organs secondary to complications associated with widespread inflammatory changes of the connective tissues. A subset of SLE, lupus profundus or panniculitis, affects approximately 2-3% of SLE patients and causes an inflammatory reaction of the panniculus, or subcutaneous adipose tissue. First described in 1971 by Tuffanelli, lupus mastitis is the clinical correlate of lupus panniculitis within the breast tissue and usually, but not always, presents with a single tender breast mass in a previously diagnosed SLE patient. Similar to most autoimmune diseases, lupus mastitis predominately affects women of childbearing age, with 37 years as the mean age at presentation. The overlying skin can be asymptomatic, or may show signs of erythema, ulceration, hyperkeratosis or poikilodermic changes[1]. Although, lupus mastitis can look and feel similar to a neoplastic tumor, this disease has a benign course. Management includes medical treatment to subdue the amount of inflammation and avoidance of surgery and other invasive procedures in order to prevent aggravation of the underlying inflammation. The physician may need to acquire imaging or tissue samples of the mass in order to confirm the diagnosis, however, a past history of SLE and a high level of clinical suspicion is often needed in order to include this rare breast finding on a differential diagnosis of a breast mass[2].

Tissue findings prove to be the most definitive way to confirm a diagnosis of lupus mastitis. Lymphocytic infiltration, occasional germinal centers and marked hyaline fat necrosis with sclerosis are considered pathognomonic findings. Lupus mastitis is a chronic inflammatory disease with remissions and recurrences being a common presentation. First line therapy includes oral hydroxychloroquine, an anti-malarial agent, with remission usually occurring in 3-6 months. More complicated cases may also require treatment with systemic steroids for an overall improvement in the patient’s symptoms. Cyclophosphamide has shown varying degrees of effectiveness as a steroid sparing treatment option. In severe cases, mastectomy may be indicated for symptom control[4].

References