Primary Synchronous Bilateral Mucosa-Associated Lymphoid Tissue Lymphoma (MALT) of the Breast: A case Report with Discussion of Management and Review of the Literature

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Abstract

Mucosa associated lymphoid tissue (MALT) lymphoma is a lymphoid neoplasm arising in extranodal sites with histologic features similar to those of nodal marginal zone lymphoma. Among the sites where a primary MALT lymphoma arises, the GI tract is the most common whereas the breast represents only 4%. Bilateral involvement of the breasts by a primary MALT lymphoma is exceptional with only 10 cases reported in the literature up to date. Herein, a case of a 73 years old woman with bilateral breast involvement by MALT lymphoma is presented and discussed.

Introduction

Mucosa associated lymphoid tissue (MALT) lymphoma is a lymphoproliferative disease arising in extranodal sites with histologic features similar to those of nodal marginal zone lymphoma. The most common site of origin of a MALT lymphoma is the gastrointestinal tract with the stomach being the most frequently involved site [1]. MALT lymphoma can arise from native mucosa-associated lymphoid tissue, such as tonsils or ileum, or from MALT that has been acquired as a result of a chronic inflammatory input or an autoimmune disease [2]. Organs where MALT is not usually present are GI tract, Lung, Salivary glands, Ocular adnexa and Breast [3].

Case Report

A 73 years old woman, with a history of uterine adenocarcinoma, hypertension and ocular myasthenia gravis, was found to have an indeterminate nodule in each breast (B-RADS-Breast Imaging-Reporting and Data System category 4) on bilateral mammograms and ultrasound. No other lesions or masses were noted in the rest of the body according to a pretreatment work-up consisting of Positron emission tomography-computed tomography (PET/CT) and blood work. Ultrasound-guided needle core biopsies of both right and left breasts showed a proliferation of atypical lymphocytes (Figure 1). Immunostains were initially performed on the left core biopsy (Figures 2 and 3) and the lymphocytic cell population was positive for CD20, CD79a (subset) and Bcl2, negative for CD10, CD23, Bcl6 and cyclinD1. CD21 was positive in follicular dendritic cells. CD138, kappa and lambda stains showed rare polyclonal plasma cells. CD3 and CD5 highlighted background T cells. Histological and immunohistochemical features were highly suspicious for a low grade B cell neoplasm, favoring extranodal marginal zone lymphoma. Fluorescence in situ hybridization (FISH) analysis for Immunoglobulin heavy locus (IgH), which is commonly associated with lymphoid disorders, was performed and the result was negative for IgH gene rearrangement. Subsequently the patient underwent right and left breast lumpectomy. Although grossly the specimens did not show any discrete nodule or masses, the histological analysis showed an atypical lymphoid cell proliferation with features identical to those present in the previous core biopsies. The morphologic changes were consistent with the diagnosis of extranodal marginal zone B cell lymphoma. Additionally the neoplastic cells resulted positive for B-cell gene rearrangement by Polymerase chain reaction (PCR), further supporting the morphologic impression. The clinical staging bilaterally was I-EA. The patient received a course of definitive radiation treatment to the right breast and lower axilla and to the left breast and lower axilla, using high tangent fields bilaterally to a dose of 39.6 Gy. The patient tolerated the course of bilateral breast radiation treatment without complication. PET CT imaging and patient’s physical examination do not show any residual disease at the most recent follow up, 1 year and 2 months after treatment.

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Among the most common sites involved by MALT lymphoma, breast represents only 4% [10]. Martinelli et al. analyzed 264 cases of PBLs from 278 patients enrolled in the International Extranodal Lymphoma Study Group (IELSG). 204 (77%) cases were DLBCL, 36 (13%) were Follicular Lymphomas, while 24 (10%) resulted to be extranodal Marginal Zone Lymphomas (MZL aka MALT). Only 1 patient had a bilateral MZL [11]. 5 of the 24 patients with MZL were treated with radiation, 5 with surgery and 1 with chemotherapy. Surgery, in association with chemotherapy or radiation or both chemo and radiation, was performed on 13 patients with MZL-PBL. In this study, the majority of patients with MZL received radiotherapy as part of the initial treatment, which was usually delivered after surgery or biopsy. Most of the relapses occurred in distant sites, with only 21% of the recurrences arising in the primary disease sites (13%) or in the contralateral breast (8%). No patients who received radiotherapy relapsed within the irradiated fields, confirming the role of this modality treatment to prevent local recurrence. Bilateral breast involvement by MALT lymphoma is rare, with only ten cases reported in the literature up to date [11-21]. A review of the literature points out that the majority of patients, with early stage unilateral or bilateral MALT breast lymphoma, has been treated with radiotherapy alone with an optimal outcome.

MALT lymphomas, regardless of stage, are among the most indolent lymphomas [3]. The transformation from marginal zone mucosa-associated lymphoid tissue (MALT) lymphoma to a more aggressive lymphoma is a rare occurrence [12]. Radiotherapy has shown excellent outcome in patients with early stage MALT [13]. Surgical options such as full mastectomy or wide local excision do not appear to offer survival benefit [14]. Therefore, radiation alone is recommended in patients with early stage PBL [9]. For patients with bilateral breast MALT lymphomas, the therapeutic management largely depends on clinical stage. There are several treatment options: radiation therapy, systemic therapy or no further therapy (watchful waiting). For selected patients with early disease presenting in the breast, radiation therapy appears to be an effective treatment option, according to NCCN guidelines® (Version 4.2014) [22].

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