Case Report

A Rare Case of Splenic Marginal Zone B-Cell Lymphoma Mimicking Relapsing Polychondritis of the Ear

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Received 1 September 2014; Revised 4 November 2014; Accepted 10 November 2014; Published 2 December 2014

1. Introduction

Auricular inflammation has a broad differential, including atopic dermatitis, contact dermatitis, cellulitis, psoriasis, systemic lupus erythematosus (SLE), other autoimmune cartilaginous conditions, chondrodermatitis nodularis, malignancy, trauma, or relapsing polychondritis (RPC). Diagnosis is typically rendered by clinical signs and symptoms but can require biopsy for a definitive tissue diagnosis.

Relapsing polychondritis (RPC) is a poorly understood phenomenon associated with cartilaginous inflammation of the ear, nose, tracheobronchial tree, and peripheral joints. Many cases of RPC respond to anti-inflammatories and resolve with no further complications. However, RPC has also been linked to more insidious conditions such as malignancies, autoimmune disorders, vasculitis, or underlying infections. Given the spectrum of associated disorders, patients with RPC may need to be monitored for more insidious underlying conditions. In this case, we report a unique case of bilateral auricular inflammation and nasal inflammation mimicking RPC as the only presenting symptom of splenic marginal zone B-cell lymphoma and we survey related cases in the literature.

Malignancy involving the ear auricles also presents as inflammation but rarely is it bilateral. We present an unusual case of RPC-like symptoms that were diagnosed as splenic marginal zone lymphoma (SMZL) with cutaneous manifestations. SMZL is a neoplasm of small B-lymphocytes that replaces white pulp germinal centers in the spleen. SMZL is relatively rare, constituting less than 1% of all non-Hodgkins lymphomas [5]. It occurs almost exclusively in patients over 50 years of age, with median age of presentation at 65 years. Patients with SMZL typically present with splenomegaly,
lymphocytosis, and cytopenia due to hypersplenism [6, 7]. SMZL does not typically have lymphadenopathy, systemic symptoms, constitutional “B” symptoms, or extra-lymphatic involvement. The incidence is twice as high in patients of Caucasian ancestry as other races, with no gender predominance.

The prognosis is generally excellent, with median overall survival in excess of 10 years. However, there is a subset of SMZL that is extremely aggressive with a median survival of 18 months [8]. Treatment is controversial, as marginal zone lymphomas are relatively rare and there are few randomized trials comparing treatments.

Given the retrospective nature of this study, it was granted an exemption in writing by the Institutional Review Board (IRB) committee of OSF Saint Anthony Medical Center.

2. Case Report

A 71-year-old Caucasian male initially presented to his primary care provider with nontender erythema of his right ear. The patient denied any trauma, fever, chills, diaphoresis, hearing involvement, weight loss, or otorrhea. Past medical history was significant for aortic and mitral valve replacements, atrial fibrillation treated with Coumadin, and chronic splenomegaly. The patient was placed on a 20 mg prednisone taper, which failed to resolve the erythema or rash after 2 weeks. In fact, there was found to be bilateral ear involvement, nasal involvement, and new onset tenderness at the 2-week follow-up. Patient was placed on 0.1% Triamcinolone EX CREA for suspected polychondritis with orders for erythrocyte sedimentation rate (ESR), anti-nuclear antibody (ANA), and rheumatoid factor tests. ESR was significant at 101 mm/h (normal 0–15 mm/h), ANA was negative at a 1:80 dilution, and RF QT was negative at <15 IU/mL.

Patient was referred to otolaryngology for evaluation. Physical examination was significant for erythematous, edematous, tender auricles bilaterally (Figures 1 and 2), and nasal tip with unremarkable findings on the rest of the exam. The left auricle had the classic lobule-sparing inflammation of RPC, whereas the right auricle presented with tenderness of the entire ear. A punch biopsy was performed due to clinical suspicion for relapsing polychondritis.

Surgical pathology (Figures 3, 4, 5, and 6) showed atypical diffuse lymphoid infiltrate of the superficial and deep dermis favoring low grade B-cell lymphoma, marginal zone subtype. Ki-67 proliferation index showed 10% of tumor cells with positive nuclear stain. Immunohistochemistry staining was equivocal for CD20, BCL-2, and CD45. Staining was negative for CD3, CD5, CD10, CD23, CD30, and cyclin D1. Outside consultation agreed with the diagnosis. Clinical correlation by hematology and oncology indicated a diagnosis of likely primary cutaneous marginal zone lymphoma (PCMZL) with plasmacytic differentiation. Oncology ordered a colonoscopy, esophagogastroduodenoscopy, staging CT scan of the chest and pelvis, and a trial of Rituxin (Rituximab).

Complete blood count (CBC) showed a normocytic anemia (hemoglobin level of 9.8 g/dL) with white blood cell count and platelets within normal limits. CT of the chest, abdomen, and pelvis showed slightly increased splenomegaly from prior CT and minimally enlarged retroperitoneal lymph nodes. There were no other changes from a baseline CT performed 4 months prior to admission. Esophagogastroduodenoscopy was unremarkable, and biopsies showed no abnormalities. Colonoscopy showed a single nonbleeding vascular ectasia and diverticulosis with no other abnormalities. Patient was then referred to a tertiary care center for further evaluation. Of note, the patient’s auricular inflammation and tenderness resolved between oncology evaluation and referral to the tertiary care center. Lactate dehydrogenase (LDH), C-reactive peptide (CRP), bone marrow biopsies, and screens for hepatitis B, hepatitis C, and HIV were performed. LDH was elevated at 279 (range 122–222 U/L) and CRP was elevated at 63.6 mg/L (normal <8.0 mg/L). Hepatitis B, hepatitis C, and HIV screens were negative. Bone marrow biopsy showed hypercellular marrow with involvement by the lymphoma, constituting approximately 10–15% of the total marrow cellularity.

A splenectomy and liver biopsy were subsequently performed. Splenectomy specimen showed splenic marginal zone lymphoma with involvement of the spleen and hilar lymph nodes. Specimen was CD5+, CD20+, and weakly CD19+. Immunohistochemistry showed no reactivity for CD10, CD23, CD3, CD7, and CD10. Patient was assessed to not require further treatments after splenectomy and would be closely followed at the tertiary center. To our knowledge, the patient has not had a recurrence of his RPC-like symptoms.

3. Discussion

Given the undefined etiology of relapsing polychondritis and the potential connection to malignancy, providers must be
**Figure 3:** Hematoxylin and Eosin stain, light microscopy $10 \times 4$ (a), $10 \times 10$ (b).

**Figure 4:** Hematoxylin and Eosin stain, light microscopy $10 \times 40$ (a), $10 \times 40$ (b).

**Figure 5:** (a) Immunohistochemistry stain for BCL-2, light microscopy $10 \times 4$. (b) Immunohistochemistry stain for CD20, light microscopy $10 \times 4$. 
vigilant in evaluating patients, particularly in cases where RPC is refractory to initial treatment. Miller et al. [9] described the first case of malignancy presenting as relapsing polychondritis in 1974, and there have been several reports of malignancies presenting as RPC since then. RPC has a strong association with leukemia and lymphomas in particular. Fransen et al. [10] reported a case of chondrosarcoma that initially presented as costochondritis, auricular inflammation, and inflammatory polyarthritis. Bochtler et al. [11] reported a case of chronic lymphocytic leukemia that presented as inflammatory polyarthritis, ocular inflammation, and bilateral auricular chondritis. Castrejon et al. [12] reported a case of unilateral left auricular inflammation, uveitis, and erythema nodosum diagnosed as lymphoplasmonic lymphoma. Lichauco et al. [13] reported a case of MALT lymphoma presenting as bilateral auricular thickening, but also with proptosis and polyarthralgia. It is evident from the literature that the constellation of symptoms in RPC may indicate a more insidious underlying systemic disease.

The treatment of SMZL is currently unclear. Initially, splenectomy was considered the first line treatment as it improved cytopenia, lymphocytosis, and progression free survival (PFS) [14]. Updated follow-up in 2013 also confirmed the benefits of splenectomy [15]. In addition, splenectomy has palliative effects on abdominal discomfort and prolonged the length of time before remission [16]. However, the overall survival, risk of histological transformation, and risk of death did not change with splenectomy. Other studies, in fact suggest that there is no difference in outcomes with or without splenectomy [16]. Rituximab, an anti-CD20 monoclonal antibody, has efficacy and complication rates equivalent to splenectomy and is an option in patients where surgery is relatively contraindicated. Milosevic et al. [17] found that patients treated with combination splenectomy and chemotherapy improved survival and disease remission when compared to splenectomy alone. Two studies found that chemotherapy with Rituximab was an effective treatment option for SMZL, superior to splenectomy, and supported the idea that splenectomy along with chemotherapy was more efficacious in inducing remission and improving survival than splenectomy alone [18, 19]. However, given the rarity of SMZL and the paucity of robust data, providers must tailor treatments to individual patients.

This is the first ever reported case of SMZL with a presentation mimicking relapsing polychondritis. The case reinforces the idea that physicians treating relapsing polychondritis must be vigilant in searching for underlying malignancies and myelodysplastic syndromes such as B-cell lymphoma, particularly when the condition does not respond to standard treatment.

Conflict of Interests

The authors declare that there is no conflict of interests regarding the publication of this paper.

References


