Dapsone for Recalcitrant Eosinophilic Annular Erythema: A Case Report and Literature Review

Luke Wallis · Robert C. Gilson · Robert T. Gilson

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ABSTRACT

Eosinophilic annular erythema (EAE) is a rare entity of unknown etiology that is possibly related to a hypersensitivity reaction and presents as annular erythematous plaques with tissue eosinophilia. It is classified as a figurate erythema with a controversial relationship to Wells syndrome (WS) in the literature, where it is generally considered a separate entity or subset based on clinical and histopathologic differences. EAE typically presents with recurrent, erythematous, arcuate, and annular plaques on the trunk and proximal extremities. The course of the disease is often chronic, recurrent, and relapsing. Responses to treatment are variable but are typically best with systemic steroids and antimalarials. We report a patient refractory to other therapies who had a striking response to dapsone.

Keywords: Annular erythema; Dapsone; Eosinophilia; Eosinophilic; Treatment resistance; Wells syndrome

INTRODUCTION

We report a rare case of eosinophilic annular erythema (EAE) and summarize the clinical features and treatments of other cases reviewed in the literature to date.

CASE REPORT

Our patient, a 38-year-old Hispanic male with a remote history of Wegener's granulomatosis and mild chronic leukopenia, presented in February 2014 with a recalcitrant, intensely pruritic, painful, burning, widespread annular eruption (Figs. 1 and 2) affecting the neck, upper thorax, arms, and proximal thighs that had been ongoing for more than two years. The patient provided informed consent to include him in this case report.

Laboratory screening showed a normal comprehensive metabolic panel and a CBC with a chronic mild leukopenia and, in some instances, a mild peripheral eosinophilia. Screening antibodies for ANA, SSA, SSB, RPR, ANCA, hepatitis B and C, and HIV 1/2 were all negative.

Multiple skin biopsies showed a superficial and deep perivascular and interstitial lymphocytic and eosinophilic infiltrate (Figs. 3, 4, and 5), some with focal vacuolar interface changes...
and prominent pigment incontinence. There were no findings of dermal edema or mucin to suggest cutaneous lupus erythematosus. No signs of vasculitis or flame figures were noted. DIF was negative for discoid lupus and systemic lupus erythematosus, with a negative LE band test reaction observed as well as a negative in vivo ANA test.

The patient was initially temporarily steroid responsive with short courses of oral prednisone, but supplemental monthly intramuscular 60–80 mg kenalog injections were necessary to achieve any noticeable control. The dermatitis continued to flare, with recurrent relapses every 3–4 weeks following each injection. Extended patch testing was positive with carba mix but there was a doubtful reaction to Balsam of Peru (Myroxylon pereirae). Balsam of Peru is a marker for a fragrance
allergy as well as allergies to certain flavorings, including vanilla, curry, cinnamon, and cloves. When foods containing this allergen are ingested, systemic dermatitis can result. The patient did trial a Balsam-of-Peru-free diet, which involved avoiding chocolate, wine, beer, cola, and tomato-based products such as pizza sauce and salsa. However, he did not experience any significant improvement with this diet, nor any improvement when he was placed on a trial of various antihistamine combinations and topical steroids. He was diagnosed with EAE and placed on hydroxychloroquine 200 mg bid, which prompted only a partial and modest clinical response. Persistent symptomatic annular lesions caused significant pain and itching and affected his quality of life.

Further options were discussed at city-wide grand rounds, including methotrexate, cyclosporine, and dapsone. Given his history of chronic mild leukopenia, cyclosporine was added next, and was incrementally increased to a dose of 200 mg bid (5 mg/kg/day), but there was no response. Oral prednisone was continued and was required to provide relief.

After a hematology evaluation for his leukopenia, dapsone was used in place of the cyclosporine, which led to a prompt response (Fig. 6), with further relief observed on dose escalation to 150 mg daily. The majority of the erythematous annular rings and plaques on the abdomen, chest, and back resolved. While he still had extensive postinflammatory hyperpigmentation, the lesions were flat and macular. He was able to taper off his prednisone completely while maintaining the response to dapsone and hydroxychloroquine. He later relapsed and required some further intermittent oral prednisone courses. However, the response to dapsone was dramatic and, for the first time, provided relief from the severe pruritic symptoms associated with this refractory skin condition, and led to an improved quality of life.

**DISCUSSION**

**Clinical Presentation**

Patients with EAE classically present with recurrent erythematous or urticarial arcuate and annular plaques on the trunk and proximal extremities [1]. Clinically, EAE is described as presenting in one of two patterns: first as a figurate/centrifugum-type pattern reminiscent of deep erythema annulare centrifugum, or second as an urticarial/annular-type pattern [2]. It is devoid of bulla, though there is a single case report of bullous eosinophilic annular erythema with negative direct immunofluorescence and negative indirect IF as well [3].

**Histopathology**

Histologically, EAE is described as having a dense superficial and deep perivascular and/or interstitial lymphohistiocytic infiltrate with abundant eosinophils. It typically presents without “flame figures,” though they have been noted in some cases and with more longstanding lesions (polymorphism, multicenter). There are some reports of focal interface changes, but generally it is absent of mucin deposition, flame figures, vasculitis, or granulomas. Basal melanosis and pigment incontinence are also reported and may account for clinically observed dusky-toned areas of hyperpigmentation [4]. While still debatable, EAE is generally regarded as being distinct from annular erythema of infancy (AEI) and Wells syndrome (WS) (Table 1). Though it also shows tissue...
# Table 1 Differential diagnosis

| Diagnosis                          | Common clinical features                                      | Pertinent histopathologic findings                                                                 |
|------------------------------------|----------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
| **Tinea corporis**                 | Annular erythematous plaques with peripheral leading scale    | Hyphae in the stratum corneum; KOH preparation is confirmatory for this superficial fungal infection |
| **Granuloma annulare**             | Nonscaly annular to arcuate plaques with a dull erythematous color; often found on distal extremities | Pattern of either palisading granulomas or interstitial histiocytes with increased mucin           |
| **Deep form of erythema annulare centrifugum** | Erythematous plaques without scale on face, trunk, or extremities | Superficial and deep, intense perivascular lymphocytic infiltrate; no eosinophilia                 |
| **Erythema marginatum**            | Erythematous, polycyclic patches or plaques that migrate within hours; usually seen in children with rheumatic fever | Superficial perivascular neutrophilic infiltrate with a few lymphocytes and eosinophils          |
| **Erythema migrans (Lyme disease)** | Large erythematous plaque that rapidly expands                | Superficial and deep perivascular and interstitial lymphocytic mixed-cell infiltrate with some plasma cells and eosinophils; spirochetes are sometimes detectable in the upper dermis with Warthin–Starry staining |
| **Erythema gyratum repens**        | Multiple annular or gyrate erythematous scaling lesions with a characteristic woodgrain or zebra-like pattern; almost always associated with underlying malignancy | Nonspecific but shows hyperkeratosis, focal parakeratosis, moderate patchy spongiosis, and a mild perivascular lymphohistiocytic infiltrate; occasionally may see eosinophils |
| **Subacute cutaneous lupus erythematosus** | Erythematous, annular or scaly plaques, often in sun-exposed regions of the skin; 80% are ANA positive, majority are SSA positive | Lymphocytic interface dermatitis, dermal edema, mucin                                               |
| **Tumid lupus erythematosus**      | Edematous erythematous plaques with minimal epidermal change, usually on the trunk | Superficial and deep perivascular and periadnexal lymphocytic infiltrates that frequently affect the eccrine coils; dermal mucin is typical and sometimes striking; no eosinophilia; positive DIF in half of patients |
| **Annular erythema of infancy**    | Annular and figurate lesions with early age of onset (typically under 1 year of age) and spontaneous resolution | Perivascular mononuclear infiltrate with few eosinophils; negative DIF; lack of “flame figures”     |
| **Wells syndrome**                 | Tender edematous cellulitis-like nodules and plaques with annular and figurate lesions | Prominent papillary edema with diffuse and deep dermal eosinophils, often into the subcutis, and classic “flame figures” |

△ Adis
eosinophilia, erythema of infancy predominantly affects children under the age of 1 year and resolves spontaneously [2]. Wells syndrome presents as tender cellulitis-type plaques, though annular and figurate lesions can occur [2]. Many physicians, however, feel that EAE is a subset or variant of WS. In WS, the inflammatory cells, including eosinophils, often involve the dermis and subcutis, whereas in EAE they are generally confined to the dermis [4]. WS also features flame figures that are quite characteristic as well as blood eosinophilia, both of which are not typical of EAE. In addition to WS, the differential diagnosis may also include the deep form of erythema annulare centrifugum, tumid lupus erythematosus, Jessner lymphocytic infiltrate, urticarial bullous pemphigoid, erythema migrans, granuloma annulare, and interstitial granulomatous dermatitis [5].

**Table 1 continued**

| Diagnosis                  | Common clinical features                                                                 | Pertinent histopathologic findings                                                                 |
|----------------------------|------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------|
| Jessner lymphocytic infiltrate | Nonscaly red patches and lumps on the face, neck, and upper back | Dense perivascular and periadnexal lymphocytic infiltration in the reticular dermis; no tissue eosinophilia; lacks interface changes, dermal mucin, and direct immunofluorescence (DIF) is negative |
| Urticarial phase of bullous pemphigoid | Large erythematous urticarial plaques, often with an annular appearance, which can subsequently convert to bullous eruptions | Eosinophilic spongiosis, upper dermal inflammatory infiltrate predominantly eosinophils; immunofluorescence shows linear IgG and C3 at the dermal–epidermal junction, as in classic bullous pemphigoid |

**Pathogenesis and Associated Diseases**

The etiology of EAE remains largely unknown, though it is suggested that it may be a hypersensitivity reaction to an unidentified antigen [6]. It has been associated with several chronic conditions, such as autoimmune thyroid disease (both hypothyroidism and hyperthyroidism), chronic borreliosis, chronic *H. pylori* gastritis, chronic kidney disease, diabetes mellitus, and hepatitis C virus infection [5, 7]. Though suspected to be a hypersensitivity response, it has also been associated with internal malignancy. Specific malignancies have been reported in three cases, including thymoma with complete resolution following thymectomy [7], metastatic prostate adenocarcinoma [8], and renal carcinoma [9].

**Treatment**

EAE is most commonly reported to respond successfully to systemic steroids and antimalarials, but it typically runs a chronic relapsing and remitting course, with a relapse frequently seen after discontinuing both treatments. It often shows resistance to multiple treatments. Most reports in the literature note the use of antimalarials as the first line for EAE. Response to hydroxychloroquine is typically prompt and usually observed within the first 4 weeks, though a case showing a delayed response with remission at 10 weeks has been reported [10]. Though Wells syndrome has been successfully treated with dapsone, there has been only one report prior to this one of a successful response of EAE to dapsone. That case report noted a complete response to dapsone at 25 mg qd, with the response occurring within 3 weeks and remission sustained at 8 months of follow-up [11]. Another multicenter study of 10 patients with EAE noted a chronic course and resistance to treatment with systemic steroids alone or in combination with hydroxychloroquine and
cyclosporine [9]. Researchers reported that EAE is characterized by a chronic course, resistance to treatment, and a high relapse rate [9]. A single case report noted a response to eight sessions of narrow-band UVB following the failure of systemic steroids and hydroxychloroquine; the light therapy cleared the EAE without inducing pigment changes, and no recurrence was noted 9 months after the light therapy had been stopped [1]. There is also a single case report of a dramatic improvement with indomethacin, but repeated syncopal episodes apparently led to the cessation of that therapy [12]. A recent case showed remission with nicotinamide at 900 mg/day dosing [13]. Though the patient had previously required systemic steroids for 7 years, remission was maintained with nicotinamide for a full year after cessation of the steroids. The patient also had a history of autoimmune pancreatitis with elevated immunoglobulin G4 levels (335 mg/dl), but the case report did not comment on the treatment of that or its timing in relation to the EAE.

As one can see from these reports in the literature, EAE commonly runs a chronic relapsing course and is resistant to numerous treatments. Though EAE typically is refractory, it can also be self-limited; treatment may not be required in all instances [6].

CONCLUSIONS

Given the rarity of this condition and the lack of a complete response to any particular therapy, dapsone should be considered as a therapeutic option for this often refractory skin condition.

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