Angiolymphoid Hyperplasia with Eosinophilia Successfully Treated with Cryotherapy

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Abstract

BACKGROUND: Angiolymphoid hyperplasia with eosinophilia is an uncommon, benign, vasoproliferative cutaneous neoplasm with uncertain origin. It preferably affects middle-aged adults, manifesting as plum-colored pruritic papules, nodules and plaques, which can persist indefinitely, relapsing over time. Different response/resistance to various therapeutic modalities and frequent recurrences impose a great therapeutic dilemma.

CASE PRESENTATION: Herein, we present a 77-year-old male patient with a 7-month-history of unrecognized cutaneous manifestations on his left shoulder and flank. Based on the investigations, the diagnosis of angiolymphoid hyperplasia with eosinophilia was established. We applied cryotherapy as a treatment of choice. The complete regression of the skin lesions and three years disease-free period was achieved.

CONCLUSION: Although surgery is the standard therapeutic approach, the disease recurs despite multiple surgical attempts. Therefore, we recommend cryotherapy as effective and safe treatment modalities for angiolymphoid hyperplasia with eosinophilia. Knowing the recurring nature of this disease, the patients with angiolymphoid hyperplasia should stay on short-term follow up in order to monitor if new lesions occur.
Case Presentation

A 77-year-old man was referred to our clinic due to 7-months history of multiple, pruritic, occasionally painful, erythematous/violaceous papulo/nodular and plaques-like lesions on left shoulder and trunk/flank (Figure 1). During that period, lesions increased in number and size, without any signs of spontaneous remission. The patient noticed crusting and bleeding from his certain lesions. He has unremarkable medical and family history. No systemic symptoms or palpable regional lymphadenopathy were detected. A laboratory workup was conducted in addition to an excisional biopsy of one skin lesion. All laboratory investigations, including serum IgE level, were within normal range.

Skin biopsy from the lesion showed prominent vascular proliferation, epithelioid-like endothelial cells, with typical presentation of "hobnail" in the lumen of the vessels (Figure 2a and 2b). Polymorphous inflammatory infiltrate composed primarily of histiocytes and eosinophils with secondary lymphoid follicles formation were also noted (Figure 3). Immunohistochemical studies showed positive CD20, reflecting the B-cell nature of the infiltrate (Figure 4). According to these findings, the confident diagnosis of angiolymphoid hyperplasia with eosinophilia was established.

Our treatment of choice was open spray techniques of cryotherapy with schedule of one session in every three weeks. Two freeze-thaw cycles with 15-s freezing and 1-min thawing per session. Number of treatment sessions was 3. A complete regression of skin lesions, without any recurrence during three years of follow-up was achieved (Figure 5).

Discussion

Angiolymphoid hyperplasia with eosinophilia has been reported worldwide with an undetermined frequency. The pathogenesis of the disease is unclear [5]. Some authors consider ALHE as a benign neoplastic proliferation of endothelial cells [11], [12]. Others proposed a theory of unusual reactive hyperplasia of vascular tissue as a response to insect bite, trauma or infection [2]. Eosinophils could be actively involved in the pathogenesis of inflammatory reaction by production of nitric oxide and eosinophilic cationic protein [13].

Additional hypothesis suggest that arteriovenous shunts may play a role in the pathogenesis. In favor of the last theory, most cases of ALHE histologically show damaged and/or tortuous arteries and veins at the base of the lesion [2]. Kempf
et al., presented evidence that ALHE or its subset may represent a T-cell lymphoproliferative disorder of low-grade malignancy [14]. This hypothesis is based on the predominance of T lymphocytes and a rearrangement in TCR receptor in some cases [14]. Therefore, immunohistochemistry could be useful in some cases. Immunohistochemistry in our patient showed CD20+ cells, reflecting the B-cell nature of the inflammatory infiltrate in dermis.

Differential diagnosis of ALHE includes Kimura disease, cutaneous lymphoma, Kaposi sarcoma, pyogenic granuloma and cutaneous metastases [5]. Kimura disease clinically presents as large deep nodules covered by normal skin associated with adenomegaly and increased serum eosinophils and IgE [12].

In our case, the immunohistochemistry rules out all other differential diagnosis.

Diagnosis of ALHE is based on clinical, biochemical and unique histopathological findings. All these criteria were fulfilled in the presented case.

ALHE proves a great therapeutic dilemma, because there are a large variety of proposed treatments, yet there is not enough data on most of them. Many therapeutic procedures have been proposed including electrodesiccation, surgical excision, Mohs surgery, cryotherapy, topical or systemic corticosteroids, topical tacrolimus, imiquimod or laser therapy [11]. Surgery may be efficient in limited lesions, but recurrences are observed in 40% of the cases, due to its difficulty to determine the surgical margins [4]. Our patient actually refused surgical procedure and opted for cryotherapy, considering our explanation about the recurring nature of ALHE. However, our therapeutic approach with cryotherapy resulted in a complete regression of the skin lesions without any recurrence in the three years follow-up duration. There is no report available in the literature to date which shows long remission period after successful treatment of ALHE with cryotherapy.

In conclusion, the pathogenesis of ALHE still remains unclear and there is no consensus on the best treatment choice. Although the disease is not life threatening, it usually presents with disfiguring lesions that affect the quality of life. Further research is needed to find an effective cure and standardized therapeutic approach for this dysmorphic and recurrent condition.

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796