Cutaneous B-Cell Pseudolymphoma Successfully Treated with Triamcinolone Acetonide

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Abstract

BACKGROUND: Cutaneous pseudolymphoma (PSL) is a reactive polyclonal benign lymphoproliferative process in the skin that simulate cutaneous lymphomas clinically, histologically, or both, predominantly composed of either B-cells or T-cells, localized or disseminated. PSL clinically manifests as solitary nodules or plaque on the face. In cases where cutaneous PSL is suspected, the most crucial part is diagnosis, to differentiate benign or malignant lesion. Diagnosis required a combination of clinical, histopathological, and immunohistochemistry examination.

CASE REPORT: A 59-year-old man presented with asymptomatic erythematous plaque on her cheek for 6 months before. Histopathological examination revealed dense small lymphocytic infiltration forming lymphoid follicles with centrum germinativum that partially destructed skin appendice glands. Immunohistochemistry examination showed positive result on cluster of differentiation (CD)20 and CD3 staining. With domination of CD20 treatment: Patient was treated with intraleisional injection of triamcinolone acetonide 10 mg/ml and showed satisfying result after 3 times injection.

CONCLUSION: A cutaneous B-cell PSL in a 59-year-old man was diagnosed based on history and physical, histopathological, and also immunohistochemistry examination. Intraleional injection of 10 mg/ml triamcinolone acetonide gave satisfying result.

Introduction

Cutaneous pseudolymphoma (PSL) is a term used for benign reactive lymphoproliferative skin disorders that mimic cutaneous lymphoma, both clinically and/or histopathologically. These diseases differ in clinical, histological, immunophenotypic presentation, and etiology [1], [2]. The term PSL was first introduced by Kaposi in 1891 and the term cutaneous PSL was coined by Burg et al. in 1982 [3].

Based on the dominant lymphocyte infiltrate, cutaneous PSL is classified into B-cell PSL (B-PSL), T-cell PSL (T-PSL), and mix PSL [4]. Cutaneous B-PSL often presents as nodeule or plaque. Immunohistochemical staining shows that most infiltrates are more dominantly stained with B-cell markers than T-cell markers. Cells in reactive follicles express BCL-6 (+) and BCL-2 (−) [1].

Cutaneous PSL may affect all age group; Borrelia-induced B-PSL more commonly occurs in children and young adults while drug-induced T-PSL is more commonly encountered in adults. Even though Borrelia-induced PSL may be a precursor for B-cell neoplasms of the skin, in general, cutaneous PSL is self-limiting and does not affect survival [2].

Clinical features of cutaneous PSL include papules, plaques, or purplish erythema nodules, which are mostly found on the face [4], [5], [6]. A broad spectrum of known causative factors that may induce cutaneous PSL has been identified. Clinicopathological correlation is central in establishing the diagnosis of cutaneous PSL and distinguishing it from skin lymphoma. Infectious agents, such as spirochetal bacteria (Borrelia burgdorferi sp. and Treponema pallidum), viruses (e.g.: Parapoxvirus), and infestations (e.g. scabies), insect bites, injection of vaccines, or desensitization antigens, foreign bodies such as tattoos and metals, and drugs have been identified as causative factors for PSL. All cases without identifiable causes are called idiopathic PSL [1], [5], [6], [7].

Here, we report one case of cutaneous B-PSL in a 59-year-old man who was successfully treated with intralesional triamcinolone acetonide injection.

Case Report

A 59-year-old man came to the Dermatovenerology Outpatient Clinic of Hasanuddin University Hospital, Makassar, Indonesia, with a chief complaint of redness lump...
on the left cheek that appeared 6 months before admission. The lump was originally in the form of skin-colored small nodules resembling acne that reddened, increased in number, and widened. The lumps were not painful nor itchy. The patient denied history of previous medication, insect bites, or trauma that preceded the complaint. Family history with similar complaints and history of diabetes were denied. History of food allergies and drug allergies was also absent.

Physical examination showed that vital signs were within normal limit. Lymph node enlargement was absent. The results of laboratory tests of complete blood count, hematologic, and blood chemistry (glucose, kidney function, and liver function) were within normal limits.

Dermatological examination revealed erythematous plaque with irregular edges on the left preauricular region (Figure 1a). On palpation, the plaque was compressible and fixed without any tenderness. The differential diagnosis included sarcoidosis, lupus vulgaris, leprosy, cutaneous PSL, and cutaneous lymphoma.

Biopsy examination was done to confirm the cause of the lesion. Histopathological examination showed normal epidermis and clusters of lymphocytes forming nodules in the dermis with some surrounded the adnexal structures (Figure 2). No atypical lymphocytes nor mitotic activity was found. Immunohistochemical staining showed cluster of differentiation (CD)3 and CD20 expression on the lymphocyte membrane, with CD20 being more dominant (Figure 3 and 4). These findings were consistent with that of cutaneous B-PSL.

Figure 1: Left preauricular region. (a) Erythematous plaque with irregular edges on the first admission. (b) Thinning of the lesion 1 week after the first injection of triamcinolone acetone. (c and d) The plaques faded to brown and thinned 1 week after the second and third injection of triamcinolone acetone

Figure 2: (a-c) Histopathological examination. (a) ×4, HE staining, epidermis was still good, dermis showed distribution of lymphocyte cells arranged in groups to form nodules (nodular structure), some areas appear diffuse and compact, and appear to surround the adnexal structure. Infiltrate consists of lymphocytes small, dense small lymphocyte form lymphoid follicles with centroblasticum which partially degrades the skin's appendices, there were eosinophils and macrophages, lymphocytes do not show atypia, no mitotic activity was found. (b) ×10, HE staining. (c) ×40, HE staining, no mitotic activity was found

Based on the history, physical, and supporting examinations, the patient was diagnosed as B-PSL. Intradermal injection of 10 mg/ml triamcinolone acetone per week was administered and the lesion showed significant improvement after three courses of injections (Figure 1b and d). The erythematous nodules faded into brownish plaques (Figure 1c) and no side effects were observed.

Discussion

In cases of suspected cutaneous PSL, excluding the possibility of malignancy is of utmost
importance [4], [8]. The diagnosis must always be made based on the combination of clinical and histopathological examination and often requires additional examinations such as immunohistochemistry and molecular biological examination [7], [9].

Clinically, most PSL cases manifest as solitary erythematous to brown or purplish plaque with facial predilection; however, it can also appear as solitary to multiple papules or infiltrated plaques [4]. Some cases also reported unusual forms in the form of ulcerations and keloids [10], [11].

Based on the clinical and/or histological presentation, there are four main groups of cutaneous PSL: Nodular PSL, pseudomycosis fungoides, other PSL (representing different clinical entities), and intravascular PSL [12], [13]. In this case, dermatological examination revealed erythematous plaque with irregular edges on the left preauricular region which was consistent with nodular PSL.

Pathologically, cutaneous PSL is classified into ordinary PSL (O-PSL), PSL with dominant B-cell infiltrates (B-PSL), PSL with dominant T-cell infiltrates (T-PSL), and PSL with mixed and unclassified infiltrate [2], [7]. In immunohistochemical examination, most infiltrates are represented by more or dominant B-cell markers than T-cell markers [1].

Antigens in cutaneous PSL can be in the form of insect bites, tattoos, trauma, vaccinations, jewelry, drugs, and infection by B. burgdorferi. In many cases, the etiology of PSL cannot be unveiled and, therefore, can be classified as an idiopathic case [7], [8], [9].

In this case, no abnormalities were found in the epidermis and positive B-cell markers (CD20) as well as a rare cell population expressing T-cell markers (CD3) were shown by immunohistochemical staining. In cutaneous B-PSL, immunohistochemical study will usually reveal the predominance of B-cells with variable numbers of T-cells. Thus, a diagnosis of B-PSL could be established [1], [14]. Important histopathological feature of PSL is lymphoid proliferation with involvement of the centrum germinativum, infiltrate consisting of small lymphocytes, a group of small dense lymphocytes forming lymphoid follicles with germinativum centrum which partially degrades the appendices of the skin, and a minimal degree of atypia. Immunohistochemistry examination typically gives positive B-cell and T-cell marker, the bias can be seen in the presence of eosinophil, macrophages, histiocytes, and dendritic cells [4], [6], [9]. The involvement of centrum germinativum and polyclonality of lymphocytes will rule out a differential diagnosis of cutaneous lymphoma [4].

Treatment options for cutaneous PSL are determined based on the causative factor, area of the lesion, anatomic location, and patient’s needs [10]. Spontaneous resolution is reported in some cases with cessation of the underlying etiology and those of idiopathic nature [10], [11]. Treatment options involve intralesional and systemic steroids, cryosurgery,
alpha interferon, excision, local radiotherapy, and immunosuppressants [4], [5], [11]. In this case, as the lesion was solitary and localized, we decided to administer intralesional triamcinolone acetonide injection. Triamcinolone acetonide injection resulted in a significant improvement after the third injection and no side effects were found in the patient. In other reports, treatment with topical and/or intralesional corticosteroids resulted in improvement in the skin disease [3]. Lesions of PSL were treated with intralesional triamcinolone acetonide injection 10 mg/ml for 2 sittings, 3 weeks apart following which resulted in complete remission of the nodules without any relapse on follow-up [7].

It is important to understand that neither clinical nor histological features alone are sufficient to distinguish PSL from lymphoma. A combination of clinical signs, histological features, and the course of the disease is needed to obtain the correct diagnosis. Sometimes, a careful drug history, serological tests, and patch tests may help to distinguish CPL from lymphoma. Nevertheless, since progression of CPL to malignant lymphoma can occur, perhaps induced by persistent antigenic stimulation, regular follow-up of the patient is mandatory. However, progression to cutaneous lymphoma has been observed in only a minority of cases [3].

**Conclusion**

Cutaneous PSL is a term used for benign reactive lymphoproliferative skin disorders that mimic cutaneous lymphoma, both clinically and/or histopathologically. Therapy with triamcinolone acetonide injection of 10 mg/ml resulted in a satisfactory outcome.

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