Palisaded neutrophilic and granulomatous dermatitis – cutaneous manifestation of Lyme disease or connected with CTD? Case report

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Abstract

Introduction. Palisaded neutrophilic and granulomatous dermatitis (PNGD) are terms which include such diseases as rheumatoid nodules, Churg-Strauss granuloma, and interstitial granulomatous dermatitis with arthritis. This heterogeneous group was first described in 1965. It is associated with immunological diseases. There are reported cases of coexistence with systemic lupus erythematosus, rheumatoid arthritis, Wegener’s granuloma, inflammatory bowel disease, generalized vascular inflammation, and lymphoproliferative disorders. The etiology of the disease is unknown. It is probable that the deposition of immune complexes in blood vessels leads to cutaneous leukocytoclastic vasculitis, degeneration of collagen fibres and palisaded granulomatous inflammation. Ultimately, this leads to fibrosis of the skin.

Objective. The aim of the study is to present a patient with skin lesions and histopathological features of palisaded neutrophilic and granulomatous dermatitis during the course of infection Borrelia burgdorferi. Several cases of this disease have been reported worldwide (30 entries in the PubMed database). To-date, there have been no reports of PNGD in Polish literature.

Materials and method. The patient, aged 72, was admitted to hospital because of erythematous, indurated lesion of the skin on the side surface of the left thigh. Diagnosis of PNGD was made on the basis of typical histopathologic features due to clinical symptoms.

Conclusions. Clinical diagnosis of PNGD is difficult, and is based mainly on the histopathological picture. Systemic therapy is incorporated mainly due to the systemic disease. The patient requires further observation in the direction of associated systemic disorders.

Key words
Palisaded neutrophilic and granulomatous dermatitis, borreliosis (Lyme disease), lupus erythematosus

INTRODUCTION

Palisaded neutrophilic and granulomatous dermatitis (PNGD) is a rare entity that has not been clearly defined, neither histopathologically nor clinically [1]. The first cases of PNGD were described in 1965 by Dykman et al., who presented two patients with skin lesions on the lateral trunk that resembled indurated linear and arcuate cords or bands, associated with rheumatoid arthritis [2]. Histopathological examination revealed an interstitial granulomatous dermatitis with lymphocytes and neutrophils, as well as areas of degenerated collagen. Nowadays, PNGD is an umbrella term used to encompass non-specific skin lesions due to such disorders as rheumatoid papules, interstitial granulomatous dermatitis with arthritis (Ackerman syndrome), and Churg-Strauss disease [3]. A PubMed search identified only 21 publications about palisaded neutrophilic and granulomatous dermatitis. No cases of this disease have been described previously in Poland.

CASE REPORT

A 71-year-old woman presented with a 4-month history of tender, smooth, erythematous plaques on the lower extremities and trunk. Physical examination revealed multiple ‘woody hard’ infiltrative lesions, erythematous-to-violaceous in colour. Most of the lesions were medium in size, measuring approximately 3–5 cm in diameter. The first lesion appeared on the left hip; surface of this plaque was rough and covered with exfoliating epidermis. A similar lesion was localized above the left knee (Fig. 1). On the trunk, there were many smaller lesions, likewise hard and firm, but the skin above the lesions was not so inflamed as the previous one (Fig. 2). The patient had no lymphadenopathy.

The eruptions did not disappeared spontaneously and new lesions appeared intermittently. She complained of diffuse myalgia and occasional fatigue with a slight fever of 37.5–37.8 °C. Review of systems including morning stiffness of phalangeal joints. No history of oral ulcers or alopecia was elicited.

Past medical history included myocardial ischaemia, hypertension, diabetes mellitus type 2 and hypercholesterolaemia.

Laboratory examinations showed sightnormochromic normocytic anaemia (Hb 10.92 g/dL, Ht 32.02%, MCV
88.6 fl, MCH 30.2 pg) with signs of inflammation (ASR 23/56mm qand CRP 12.4 mg/L). Antinuclear antibody titer was 1:320 with a homogenous pattern. Anti-Ro/La, anti-double-stranded DNA, anti-cardiolipin, anti-CCP, and anti-SCL-70 antibodies were negative. A complete blood count, basic metabolic panel, hepatic function tests, iron level tests (TIBC, UIBC, ferritin and transferrin), C3 and C4 levels, and urinalysis were normal. Hepatitis B antigen, C antibody and tumour markers detected by immunohistochemistry were negative. Serological tests for Lyme disease revealed positive reactions in the serum for IgG antibodies (>30 BBU/ml) and negative for IgM antibodies (0.7 BBU/ml). Photo test (exposure to UVB) was positive. Minimal dose that caused erythema in 24 hours was 0.038J/cm2. Punch biopsy was performed to aid the diagnosis of the eruption. Within the superficial and deep dermis there was a perivascular interstitial mixed-cell infiltrate, composed of neutrophils, histiocytes and lymphocytes. In some foci, the inflammatory infiltrate was palisaded and associated with basophilic alteration of collagen (Fig. 3). These findings were interpreted as PNGD. Narrow linear IgG and C3c staining of the basement membrane was seen in direct immunofluorescence (Fig. 4). The patient was treated with cefuroxin 2 x 500mg p.o., topical methyprednizolon and aceponas was also prescribed.

DISCUSSION

Chu et al. defined palisaded neutrophilic granulomatous dermatitis in 1994 as a disease in which circulating immune complexes occur and can develop a papular eruption on the extremities [4]. The authors suggest that PNGD represents a continuum with an inciting lesion representing leukocytoclastic vasculitis, and with palisaded granulomas observed in fully developed lesions [5]. The term ‘palisaded’, when used in dermopathology, refers to the arrangement of histiocytes in an inflammatory infiltrate [2]. The lesions of palisaded neutrophilic and granulomatous dermatitis are usually symmetrically distributed on the extremities and have smooth, ulcerated, or umbilicated surfaces [6]. However, PNGD can present as pink to violaceous papules, plaques, nodules or linear bands, some of which can have urticarial or annular configurations [5]. Early lesions are urticaria-like annular plaques, or may have a livedoid appearance. In later stages, the lesions are more infiltrative and pleomorphic (e.g. violaceous annular plaques, waxy papules, painful subcutaneous nodules, indurated linear bands) [5].

Finally, in the presented case, fibrosis was observed. The overlying epidermis is usually normal, although sometimes crusts and ulcer appear. Linear subcutaneous indurated...
cordlike bands, coined the ‘rope sign’, first described in 1993 by Ackerman, are pathognomonic of interstitial granulomatous dermatitis [7]. The duration of the disease in the presented case was not long, but the lesions were more advanced than observed in another cases.

PNGD is heterogenous entity associated with systemus lupus erythematosus, rheumatoid arthritis, Wegener’s granulomatosis, inflammatory bowel disease, and lymphoproliferative disease [3, 8]. Single cases of Behcet disease, hepatitis, sarcoidosis, leukocytoclastic vasculitis, systemic sclerosis has been reported [9, 10, 11, 12, 13]. Recently, two groups of scientist, Newell et al. and Misago et al., reported cases of patients with PNGD with no underlying illness [11, 14, 15]. The patient in the current case presented 2 of 11 SLE symptoms (ACR): photosensitivity, positive ANA, and additionally, linear IgG and C3c staining of the basement membrane in direct immunofluorescence. She had positive IgG antibodies against Borrelia burgorferi, through which Lyme disease was recognized.

The etiology of PNGD is unknown, although in 1983, Finan and Winkelmann, using direct immunofluorescent examination, observed IgM and C3 in small vessels [16]. They presumed that the cutaneous lesions were the result of immune complexes generated by underlying systemic diseases [16]. Similar observations were seen in the presented case, there was narrow linear IgG and C3c staining of the basement membrane.

Differential diagnoses include interstitial granuloma annulare, leucocytoclastic vasculitis, urticarial vasculitis, Sweet syndrome, rheumatoid nodules, necrobiosis lipoidica, and erythema elevatum et diutunum [1, 8]. Most eruptions are asymptomatic, but our patient complained tenderness. Sometimes pruritus appear [8].

Various treatment modalities tried include oral steroids, dapsone, colchicine, cyclosporine, cyclophosphamide, hydroxychloroquine, and nonsteroidal anti-inflammatory agents, with variable success rates. Oral steroids appear to be the most beneficial therapy [1, 17]. Brecher tried to treat individual lesions with intralesional glucocorticoid injections, with some improvement [6]. Many cases have not responded to therapy and recurrences were common after treatment with prednisone was tapered [6, 7]. The condition may persist from several months to a few years [8]. Usually, the clinical course of PNGD appears to be self-limited, but this was not observed in the presented patient. Whether the appearance of PNGD in an autoimmune disease is associated with a worse therapeutic response, or a poor prognosis, has not yet been clarified [11].

Due to Al-Daraji et al., the authors of the presented case consider that the lack of silent diagnostic features, both clinically and pathologically, a high index of suspicion is mandatory for making the correct diagnosis [3].

REFERENCES


