IDIOPATHIC PYODERMA GANGRENOsum
OR A SYSTEMIC DISEASE PREDICTOR?

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ABSTRACT
Pyoderma gangrenosum (PG) is an auto inflammatory non-infectious neutrophilic dermatosis often presenting with pustules or nodules that progress to ulcers. PG has specific clinical features and non-characteristic histology findings. Some systemic diseases such as Crohn’s disease, ulcerative colitis, rheumatoid arthritis, hematologic malignancy or autoimmune disease are commonly associated with PG. One of the most characteristic features of pyoderma gangrenosum is pathergy, which is the appearance of new lesions at sites of trauma, including surgical wounds.

We present a 62-year-old male with a large ulceration on the left lower leg after a bee sting.

Keywords: pyoderma gangrenosum, neutrophilic dermatosis, ulcers, pathergy

INTRODUCTION
Pyoderma gangrenosum (PG) is a rare inflammatory neutrophilic dermatosis of unknown etiology. It may be associated with a coexisting systemic disease, most commonly Crohn’s disease, ulcerative colitis, rheumatoid arthritis, hematologic malignancy or autoimmune disease (1, 2). In most of the cases the initial lesion is a papule, pustule, or nodule appearing after minor trauma, that progresses to painful, slowly growing ulcer. It is often misdiagnosed as a soft tissue infection that can coincidentally improve due to the systemic antibiotic therapy and wound care (1, 2, 3).

We present a case of pyoderma gangrenosum persisting for 7 months, not associated with systemic disease.

CASE REPORT
A 62-year-old male presented to a dermatology clinic in 2019 with a history of bee sting after which a non-healing painful ulcer developed on the anteromedial aspect of the left shin. The lesion was large and deep, with a diameter of 15/20 cm, elevated borders and erythematous periphery (Fig. 1, Fig. 2).

Previously, the patient was treated with local antiseptic, proteolytic enzyme and a systemic antibiotic with unsatisfactory improvement.
Idiopathic Pyoderma Gangrenosum or a Systemic Disease Predictor?

The laboratory tests showed: erythrocytes (RBC) – 3.03 10^12/l; erythroblasts – 10^9/L; hemoglobin (HGB) – 118.0 g/L; hematocrit (HCT) – 0.326 l/l; MCV – 107.7 fl; MCH – 39.0 pg; MCHC – 362.0 g/L; granulocytes (Gran) % - 77.6%; lymphocytes (Lym) % - 15.0%; iron binding capacity (IBC) – 40.3 μmol/L; gamma-glutamyl transferase (GGT) – 94.0 U/L; procalcitonin – 0.4249 ng/mL; hsCRP – 70.4 mg/L; ferritin – 365.58 ng/mL; bilirubin total – 12.9 μmol/L; bilirubin direct – 4.8 μmol/L. Urine tests: blood ++; ketone bodies (+); urobilinogen ++; bilirubin +++; leucocytes – neg. Microbiology culture test of a sample taken from the ulcer found *Pseudomonas aeruginosa*.

Doppler ultrasound of the leg, chest X-ray, gastro-, colonoscopy and abdominal ultrasound of the patient found no significant abnormalities.

A hematologist and a gastroenterologist were consulted and they recommended treatment with folic acid, vit. B6, vit. B12 and ursodeoxycholic acid.

Skin biopsy from the border of the lesion was performed. The histologic findings are: pseudoepitheliomatous hyperplasia with intraepidermal neutrophilic abscesses and spongiosis of the epidermis. In the dermis diffuse inflammatory infiltrate containing numerous neutrophils was observed (Fig. 3, Fig. 4, Fig. 5).

The clinical course with pathergic phenomenon, unresponsiveness to antibiotic therapy and histopathologic features of sterile neutrophilic infiltration led to diagnosis pyoderma gangrenosum.

Systemic therapy with ciprofloxacin 200 mg twice daily, 6-methylprednisolone 40 mg i.v., dimen-diphenyl sulfone (DDS) 50 mg twice daily and
esomeprazole was initiated. Locally, KMnO4 compresses were applied.

Clinical improvement was observed in the first 10 days of the therapeutic regimen. Significant part of the devitalized tissue was removed. The size and the depth of the ulcer decreased. There were no signs of bacterial contamination (Fig. 6).

**DISCUSSION**

Pyoderma gangrenosum (PG) is a neutrophilic dermatosis, with characteristic clinical and non-characteristic histology findings, first described by Brocq et al. in 1916 as “phagedenisme geometrique/geometric phagedenism” (2,3,4). Although the etiopathogenesis of PG is still unclear, there are increasing evidence that it shares common pathogenic mechanisms with other autoimmune disorders (5,6,7). Neutrophilic dysfunction, systemic inflammation, and associated genetic factors are all involved in the formation of PG ulcers (8,9).

Previously, there were no clinical criteria distinguishing PG from necrotizing soft tissue infections (10). Recently, E. Maverakis et al. suggested one major and 4 of 8 minor criteria that can be used to diagnose PG with high specificity (11). The major criterion is biopsy of an ulcer edge demonstrating neutrophilic infiltrate and the 8 minor criteria are: (1) exclusion of infection; (2) pathergy; (3) history of inflammatory bowel disease or inflammatory arthritis; (4) history of papule, pustule, or vesicle ulcerating within 4 days of appearing; (5) peripheral erythema, undermining border, and tenderness at ulceration site; (6) multiple ulcerations, at least 1 on an an-
Idiopathic Pyoderma Gangrenosum or a Systemic Disease Predictor?

terior lower leg; (7) cribriform or “wrinkled paper” scar(s) at healed ulcer sites; and (8) decreased ulcer size within 1 month of initiating immunosuppressive medication(s) (11). Applying those criteria yields 86% to 90% specificity and sensitivity.

The presented case showed: typical histopathology findings, pterygium, erythematous periphery of the ulcer, typical anterior shin localization and therapeutic effectiveness of the immunosuppressive treatment.

Pterygium is an important sign supporting the diagnosis of PG (12,13). It is defined as development of skin lesions that resist healing after minor tissue trauma as in our case. The lesion classically begins as a pustule, vesicle, or nodule that rapidly progresses into a painful ulcer or erosion with raised borders (14,15). That type of lesions can heal spontaneously or coincidentally with antibiotic treatment. Often patients believe they have been bitten by a spider or other type of insect (16). Early aggressive tissue debridement can worsen the therapy response due to pterygium phenomenon.

The clinical course of PG is not predictable, therefore the treatment should be individualized (17). Differential diagnosis of PG includes infectious (atypical mycobacterial ulcers, cutaneous tuberculosis, cutaneous leishmaniasis, sporotrichosis, ecthyma gangrenosum, syphilis) and noninfectious diseases (vasculitis, cutaneous malignancies, drug-induced conditions) (18). Skin biopsy and microbiology tests are crucial for the diagnosis. In the presented case differential diagnosis with ecthyma gangrenosum is important, because of the presence of Pseudomonas aeruginosa in the microbial culture test. The typical massive neutrophilic infiltrate confirms the diagnosis of PG (in contrast, the histopathology of ecthyma gangrenosum shows vascular necrosis with few inflammatory cells). We would interpret the presence of Pseudomonas aeruginosa as contamination of a preexisting PG lesion, which is common for cases of ulcers with prolonged healing.

The treatment of PG remains challenging. Thorough physical examination and comorbidity investigation are crucial for the precise diagnosis and treatment of PG. For mild cases high-potency topical steroids, intralesional application of triamcinolone acetonide and calcineurin inhibitors are mostly used with good therapeutic response (19,20). For severe PG, first-line therapy includes systemic steroids or cyclosporine (21). Second-line options are: methotrexate, azathioprine, mycophenolate mofetil, cyclophosphamide, dapsone, thalidomide, colchicine and intravenous immunoglobulins used to decrease the steroid dose in maintenance therapy or in combination with first-line agents for recalcitrant PG (21). Recently, biological agents are also used for treatment of PG (21, 22). TNF-α inhibitors prescribed for inflammatory bowel diseases (Crohn’s disease and ulcerative colitis) can also improve PG associated with these conditions (23). IL-1 is the key inflammatory mediator triggering release of chemokines involved in neutrophil recruitment and activation. Anakinra and canakinumab are both therapies that block IL-1 and have been used to treat refractory PG (24). New clinical trials show that biologic agents could be an alternative for treatment of severe and refractory PG cases (25,26).

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