Chronic Recurrent Neutrophilic Dermatosis: A Possible Variant in the Spectrum of Neutrophilic Dermatoses

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Abstract. Neutrophilic dermatoses constitute a clinically heterogeneous group of diseases that share a common histological substrate, consisting of a dense dermal inflammatory infiltrate of mature polymorphonuclear neutrophils and no evidence of vasculitis. We describe the case of a 56-year-old man with a 6-month history of painful generalized erythematous edematous plaques. Histopathology indicated neutrophilic dermatosis but the patient did not have fever, elevated white blood cell count, or systemic involvement. Tests to rule out possible inflammatory, neoplastic, or infectious processes were negative. We consider the term chronic recurrent annular neutrophilic dermatosis, first used by Christensen et al, to be the most appropriate to define this variant with clinical findings that differ from classic Sweet syndrome.

Key words: neutrophilic dermatosis, chronic disease, recurrence.
arthromyalgia and presented mild bilateral conjunctivitis. Complete blood count showed 4100 white blood cells/µL with 74% polymorphonuclear neutrophils. The erythrocyte sedimentation rate was 31 mm/h. The antinuclear antibody test was positive at low titers (1/40). Anti-DNA antibody tests were negative. Chest X-ray, abdominal ultrasound, upper gastrointestinal endoscopy and colonoscopy, and bone-marrow aspiration and biopsy showed no abnormalities. All blood tests performed to rule out associated infections, including hepatotropic virus, human immunodeficiency virus, cytomegalovirus, syphilis, and herpes simplex were negative. Histopathology of a pectoral skin biopsy indicated a dense neutrophilic inflammatory infiltrate in the upper dermis without findings suggestive of leukocytoclastic vasculitis (Figure 4). Cutaneous lesions improved after beginning treatment with oral prednisone (50 mg/d), with gradual tapering of the dose. Occasional mild recurrences were reported.

Discussion

Sweet syndrome has been associated with a wide range of processes and situations, including infectious and inflammatory diseases, neoplastic disease, drug use, vaccination, and even pregnancy.9-11 Nevertheless, these conditions can only be identified in approximately 30% of cases. An occult neoplasm can be identified in 20% of cases, mainly hematologic neoplasia (85%), the most frequent types being acute myelogenous leukemia and lymphomas.12 The absence of factors commonly associated with triggering classic Sweet syndrome (upper respiratory tract infections, drugs, etc), male predominance, presence of more severe generalized cutaneous lesions, frequent extracutaneous disorders, absence of neutrophilia, presence of anemia or...
abnormal platelet count and, in particular, the high rate of recurrence, are clinical observations that frequently appear in the context of malignancy-associated Sweet syndrome. On the other hand, in these cases, cutaneous lesions can precede the onset of neoplastic processes even by several years.

In 1989, Christensen et al described 2 cases of patients with chronic and recurrent outbreaks of generalized annular erythematous, edematous cutaneous plaques, with histopathological findings suggestive of Sweet syndrome, but without fever or general symptoms. Although some cases of recurrent neutrophilic dermatosis in isolation had been described previously, Christensen et al were the first to regard this as an entity in itself, coining the term chronic recurrent annular neutrophilic dermatosis to refer to this variant of neutrophilic dermatosis with histopathological findings typical of this entity, but with a chronic and recurrent course differing from classic Sweet syndrome and no evidence of associated systemic disease.

Subsequently, Romero et al reported 4 cases of patients with lesions clinically or histologically compatible with Sweet syndrome, but with an afebrile and recurrent course lasting several years, without presenting leukocytosis or peripheral neutrophilia at any time. They referred to these cases as chronic recurrent afebrile neutrophilic dermatosis and, unlike Christensen et al, regarded this process as a variant of Sweet syndrome and not as a separate clinical entity.

From the clinical standpoint, the absence of fever and leukocytosis is the norm in most chronic and recurrent cases, although these criteria may be present in some instances. On the other hand, some of the forms of chronic recurrent neutrophilic dermatosis described present as single lesions, especially on the face. Furthermore, this chronic variant does not seem to respond so well as classic Sweet syndrome to systemic corticosteroids, and recurrence is frequent, as in our case.

Our patient shares several of the aforementioned clinical criteria that are more often associated with Sweet syndrome. However, to date, all the tests performed to diagnose this disease have been negative. Thus, we believe that the term chronic recurrent annular neutrophilic dermatosis best describes our patient’s disease. In any case, the fact that cutaneous lesions can precede the diagnosis of malignancy by several years suggests that chronic recurrent annular neutrophilic dermatosis can be an early manifestation of systemic disease such as cancer and thus clinical follow-up in these patients is recommended.

In conclusion, we describe a case of neutrophilic dermatosis of atypical, chronic, and recurrent evolution. Further studies on a large series of similar cases may help us to understand whether we are facing a true variant within the spectrum of neutrophilic dermatosis or, on the other hand, whether the condition is a severe form of Sweet syndrome as an early manifestation of a systemic disease.

Conflicts of Interest
The authors declare no conflicts of interest.

References