Fixed cutaneous sporotrichosis

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INTRODUCTION

Sporotrichosis is a subcutaneous fungal infection first described by Benjamin Schenks in 1898. It is caused by the dimorphic fungi complex Sporothrix [1,2] as the result of traumatic inoculations with contaminated material, usually spindles and thorns. Thus, this disease is popularly known as “rose gardener’s disease”, though it may also be transmitted by cats [1,2]. It is considered as a mycosis of global distribution, but its observed with more frequency in tropical countries, having higher incidence in Latin America [1,3]. The clinical types of sporotrichosis are lymphocutaneous sporotrichosis, fixed cutaneous and extracutaneous. Moreover, the heterogeneous morphology of lesions often makes the clinical diagnosis difficult, thereby leading to delayed treatment [1-3].

CASE REPORT

Female patient, 46 years, who consulted a private doctor due to her dermatological condition in her face region was sent for diagnosis. Upon examination, a dermatosis was found on the left cheek, a plaque with ulcerations on an erythematous base (Fig. 1). The rest of the physical exam was within normal limits. Familiar and personal history was found negative.

The patient indicated that the disease began 3 months ago with a small asymptomatic “pimple” on the left cheek, discharging pus, and afterwards, slowly increased and ulcerated. She received different antibiotics and non-specific creams with no effective results. Three clinical diagnosis were made based on the clinical data: inflammatory, neoplasic or infectious disease. A biopsy was performed, and the histologic sections of skin stained with HE, showing epidermis with hyperkeratosis and irregular acanthosis. From the papillary dermis to the deep reticular dermis, a diffuse granulomatosis infiltrate present at cutting surface was observed (Fig. 2).

At a higher magnification, the infiltrate was formed by granulomas with suppurative center, lymphocytes, histiocytes, plasma cells and abundant giant cells of Langhans type. Moreover, large number of yeast was observed presenting an elongated and oval shape, which was evident on PAS staining (Fig. 3). Afterwards the clinical diagnoses of fixed cutaneous sporotrichosis was performed. Upon re-interrogation, the patient revealed receiving a blow on her left cheek with a

How to cite this article: Chang P, Vásquez Acajabón MV, Logemann H, Pacheco GC. Fixed cutaneous sporotrichosis. Our Dermatol Online. 2019;10(1):44-47.
Submission: 13.03.2018; Acceptance: 07.06.2018
DOI:10.7241/ourd.20191.10
can of gaseous water about 9 months ago. A fungus cultivation was carried in Sabouraud’s glucose agar, surface growth was brown, glabrous and yeast like, with reverse also brown (Fig. 4). The microscopy confirmed *Sporothrix schenckii* (Fig. 5), the microscopic morphology of the colony was septate, branching hyphae and cluster spores. Moreover, conidia are in a flowerette arrangement at the single conidiophores (Fig. 6). The final diagnosis of the patient was fixed cutaneous sporotrichosis and itraconazole 200 mg daily was prescribed for 6 weeks, resulting in clinical cured (Fig. 7).

Prior to the study, patient gave written consent to the examination and biopsy after having been informed about the procedure.

**DISCUSSION**

Sporotrichosis, or rose gardener’s disease [1], was first described in 1898 by Benjamin Schenk at Johns Hopkins Hospital and was named by Hektoen and Perkins in 1900 [1,2]. The disease is a polymorphic subacute or chronic subcutaneous mycosis. Sporotrichosis is dispersed worldwide and more prevalent in tropical and subtropical areas, with the highest prevalence in Central and South America, Africa and Japan [1,3,4]. Sporotrichosis is caused by a complex group of dimorphic fungus species called *Sporothrix* [1,5-7], the major species includes *Sporothrix brasiliensis*, *S. schenckii*, *S. globosa* and *S. albicans* and *S. mexicana* [8-11]. *Sporothrix* is present on the natural environment in soil, dead wood (Splinter and thorns favor the growth), mosses and cornstalks [12]. But has never been observed as plant pathogen, probably due to the antifungal activity of plants [10].

It grows in the environment in a mold-like form at temperatures between 25-30°C. Moreover, it develops
into yeast-like form in host tissues at 35-37°C [5,13]. It is considered an occupational disease of particular risk for agriculturists, gardeners, florists and foresters [2,5]. Furthermore, alcoholism and diabetes have also been correlated as risk factors associated to it [2].

This mycosis is more commonly seen in tropical and subtropical countries, principally in Mexico, Central America and Peru. However, the region of Abancay in Perú is considered hyperendemic with an incidence of 48 to 60 cases per 100 000 population [11].

This subcutaneous mycosis strongly depends on exposure [12]. It affects individuals regardless of age and gender and it is easily acquired by in children and young adults. Traumatic inoculation is the typical mode for acquisition of cutaneous infection in immunocompetent hosts [5,7], and typically affects the face region and extremities. More recently, it has also been found to be transmitted via domestic cats and rodents in Brazil and Japan [1,8,14].

Several factors such as host immunity, virulence of the inoculated strain and the depth of traumatic inoculation may determinate the different types of presentation [3].
Conventionally, three clinical forms are described: 1) Fixed Cutaneous (nodulopapular, ulcerative, verrucous and furunculoide) [10,11], 2) Lympho-Cutaneous (most commonly) and 3) Disseminated [1,5].

Fixed cutaneous variety of sporotrichosis remains localized and is less common than the lymphocutaneous type [5,6]. Clinically, initial lesion develops at the site of skin inoculation [10], commonly on the face region in children and on the arms in adults [14]. This form usually presents a single and painless infiltrated, erythematous or violaceous plaque that may become ulcerated or verrucous without lymphatic involvement [2,6]. Also it may start with an aceniform or nodular form of variable duration [5,14].

The lymphocutaneous form is the classical presentation of sporotrichosis and the systemic disease is the product of the conidia inhalation or hematogenous dissemination from primary sites. Pulmonary sporotrichosis displays a form radiographically indistinguishable from tuberculosis and histoplasmosis in patients with severe underlying chronic obstructive pulmonary disease. Osteoarticular sporotrichosis result from the direct inoculation or hematogenous dissemination, with involvement of multiple visceral organs, this occurs almost exclusively in persons with AIDS [10].

Differential diagnoses include cutaneous chronic infections such as cutaneous leishmaniasis in its localized form, but the typical ulcer is painless with erythematous base and well defined high edges [6]. It must be differentiated from tuberculosis, sarcoidosis, paracoccidioidomycosis, chromoblastomycosis, leprosy and other mycobacterial diseases [2,9]. Clinical suspicion is essential for diagnosis. Moreover, direct examination of exudate is not suitable due the structures not being observed [2]. The histopathological examination of tissue stained with conventional HE (hematoxylin & cosin) lacks sensitivity [5]. The gold standard for diagnosis is the isolation of the fungus in Sabouraud dextrose agar with chloramphenicol [2,11].

Prognosis is mainly excellent when proper pharmacological treatment is instituted. Currently, oral itraconazole has become the drug of choice in a recommended dosage of 100-200mg daily, to be administered for 3-6 months [5]. However, the choice of therapy depends upon the location and form of the disease. Potassium iodide concentrated solution in dosage of 3-6 grams daily for 3-4 months, has been described more affordable and equally effective option in cutaneous forms, especially in endemic areas of developing countries [2,3]. Another therapeutic option is terbinafine, which shows similar results to those of itraconazole [4]. Heat application to lesions may help since low temperatures are preferred by fungus [10].

CONSENT

The examination of the patient was conducted according to the Declaration of Helsinki principles.

REFERENCES