A clinico-epidemiological study of polymorphic light eruption
Lata Sharma, A. Basnet

A clinico-epidemiological study of PLE was done for a period of one year to include 220 cases of PLE of skin type between IV and VI. The manifestation of PLE was most common in housewives on sun-exposed areas. Most of the patients of PLE presented with mild symptoms and rash around neck, lower forearms and arms which was aggravated on exposure to sunlight. PLE was more prevalent in the months of March and September and the disease was recurrent in 31.36% of cases.

Comparative study of efficacy and safety of hydroxychloroquine and chloroquine in polymorphic light eruption: A randomized, double-blind, multicentric study
Anil Pareek, Uday Khopkar, S. Sacchidanand, Nitin Chandurkar, Geeta S. Naik

In a double-blind randomized, comparative multicentric study evaluating efficacy of antimalarials in polymorphic light eruption, a total of 117 patients of PLE were randomized to receive hydroxychloroquine and chloroquine tablets for a period of 2 months (initial twice daily dose was reduced to once daily after 1 month). A significant reduction in severity scores for burning, itching, and erythema was observed in patients treated with hydroxychloroquine as compared to chloroquine. Hydroxychloroquine was found to be a safe antimalarial in the dosage studied with lesser risk of ocular toxicity.
Many faces of cutaneous leishmaniasis
Arfan Ul Bari, Simeen Ber Rahman

Symptomatic cutaneous leishmaniasis is diverse in its presentation and outcome in a tropical country like Pakistan where the disease is endemic. The study describes the clinical profile and atypical presentations in 41 cases among 718 patients of cutaneous leishmaniasis. Extremity was the most common site of involvement and lupoid cutaneous leishmaniasis was the most common atypical form observed. Authors suggest that clustering of atypical cases in a geographically restricted region could possibly be due to emergence of a new parasite strain.

Forehead plaque: A cutaneous marker of CNS involvement in tuberous sclerosis
G. Raghu Rama Rao, P. V. Krishna Rao, K. V. T. Gopal, Y. Hari Kishan Kumar, B. V. Ramachandra

In a retrospective study of 15 patients of tuberous sclerosis, eight patients had central nervous system involvement. Among these 8 cases, 7 cases had forehead plaque. This small study suggests that presence of forehead plaque is significantly associated with CNS involvement.

Ligand-binding prediction for ErbB2, a key molecule in the pathogenesis of leprosy
Viroj Wiwanitkit

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INSTRUCTIONS FOR AUTHORS

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Multiple xanthogranulomas in an adult

Sir,

A 21-year-old old unmarried male presented to our Department with a 2 year history of numerous well-demarcated, firm, rubbery brownish yellow papular lesions of different sizes, over the face, neck, trunk and upper extremities, the majority being on the face. The patient was otherwise healthy. General examination of all vital systems was normal. Cutaneous examination revealed multiple skin-colored and hyperpigmented papules of different sizes, distributed over the face, neck, extensor and flexor aspects of the upper extremities [Figure 1]. The trunk had a few lesions on the front and back. The lesions were asymptomatic, firm to rubbery in consistency and non-tender. Examination of mucous membranes, palms and soles was normal. The patient was a healthy active male, cycle mechanic by profession and denied any sexual contact of any type in the past. As per the patient’s version, lesions first started 2 years back on the face as a solitary papule over the right cheek, subsequently followed by appearance of multiple lesions in different parts as mentioned. A clinical diagnosis of late onset juvenile xanthogranuloma (JXG) was made and a biopsy was done. Histopathological examination revealed histiocytic proliferation with features of secondary ‘xanthomization’ with the presence of foam cells, foreign-body giant cells and Touton giant cells, in the superficial dermis. The epidermis was thinned out and without any grenz zone. Lymphocytes, eosinophils and neutrophils were variably seen in the inflammatory infiltrate with absence of plasma cells. The histopathology confirmed the diagnosis of JXG. The patient was advised radiography of skull and chest, but results were non-contributory. Sonography of the abdomen and pelvis was also normal. Hematological and biochemical parameters including lipid profile were within normal limits. A peripheral smear examination ruled out any hematological malignancy. The patient was sent for an ophthalmologic checkup and fundoscopy but no abnormality was detected. Looking at the benign course of the disease and instances of spontaneous regression and absence of any specific therapy, the patient was discharged with counseling and advised to report for a regular yearly follow-up.

Helwig and Macknay first coined the term juvenile xanthogranuloma in 1954,[1] as a benign, asymptomatic and common self-healing disorder of non-Langerhans cell histiocytosis (LCH), affecting mostly infants, children and rarely adults. Eighty per cent cases appear in the first year of life [2,3] and 20-30% cases present at birth.[2] There is no sexual or racial predilection.[4] Clinically, in 90% of JXGs, cutaneous lesions are solitary, with the head and neck being the most common sites of involvement. Extracutaneous sites involving the eye, lung, abdominal viscera and skull have been reported by many authors. Adult JXGs rarely regress spontaneously and reports of concomitant extracutaneous lesions are rare.[5]

Juvenile xanthogranuloma is the most common form of non-X histiocytosis.[6] It forms a heterogeneous group defined by the proliferation of cells with macrophage characteristics. It is important to recognize multiple adult xanthogranulomas, because of its good prognosis and the absence of visceral involvement, therefore requiring no investigations or aggressive treatments, a very important observation made by Punithwavathy et al., in their case report of adult onset xanthogranuloma.[7] Three main clinical forms are recognized: a small nodular/papular (2-5 mm); large nodular (5-20 mm); and giant xanthogranuloma (more than 20 mm).[8] But unusual clinical variants like mixed form, subcutaneous form, JXG en plaque have been reported recently.[9] Multiple xanthogranulomas are extremely unusual in adults and that to occur in an eruptive manner is quite rare and three such cases have been associated with hematological malignancy.

Figure 1: Numerous papular lesions over the face
Clinical backgrounds of patients with adult-onset xanthogranuloma are somewhat different from those of patients with juvenile xanthogranuloma, but the histological findings of both forms of the disease are identical. Extracutaneous involvement of the eye orbit, lung, liver, testis, central nervous system, kidney etc. has been reported in childhood variants, but is not seen in the adult type. Though there has been established association of JXG with neurofibromatosis (NF-1) and juvenile chronic myelogenous leukemia (JCML) in the childhood type, this has never been reported in adults. Evaluation for extracutaneous JXG is not indicated, unless there are symptoms or findings suggesting their presence, as they also disappear spontaneously. Differential diagnosis includes molluscum contagiosum, cryptococcosis, benign cephalic histiocytosis (seen exclusively in children, infiltrate lacks foamy cells and multinucleated giant cells), generalized eruptive histiocytosis (absence of granulation and lipidation), xanthoma disseminatum (lesions tend to merge into plaques, mucous membrane involvement, associated diabetes insipidus and different biopsy findings), papular xanthoma (JXG histologically recognized by its pure primitive histiocytic phase and presence of inflammatory cells, not seen in papular xanthoma). The importance of presenting this case is to highlight the fact that while making a diagnosis of common disorders like molluscum contagiosum, one must keep in mind the adult form of xanthogranuloma in differential diagnosis.