Eruptive Pseudoangiomatosis in Adult Associated to Parvovirus B19

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Introduction

Eruptive pseudoangiomatosis (EP) is a rare disease characterized by benign spontaneous regressing exanthema of an unknown etiology. It consists of an acute onset of small eruption of distinctive erythematous angioma-like papules that are often surrounded by a pale halo [1-3]. Histologic findings show dilated dermal blood vessels with plump endothelial cells protruding into the lumen and mild-to moderate perivascular lymphohistiocytic infiltrates [4]. The etiology of this eruptive disease is still unknown. However, association with insect bites [5] and with ECHO virus, coxsackie B, Epstein-Barr virus and cytomegalovirus [6,7] have been described. We are reporting a case of a woman with this condition associated to parvovirus B19.

Case Report

A 56-year-old female patient visited our Primary Care center in July 2020. The current condition begins 24 hours prior to consult, with a generalized onset of 3-4mm erythematous papules with a peripheral pale halo. Lesions were moderate pruritic, and they were located on her upper and lower extremities and trunk mostly (Figure 1) No systemic symptoms, including fever, were present. She had no history of upper respiratory tract infection and no medical problems. We took a skin biopsy and histopathological findings showed a mild superficial perivascular dermatitis. Laboratory findings revealed an acute Parvovirus B19 (IgM+). No other blood test findings were remarkable. Treatment was started with cetirizine 10mg per day and the lesions spontaneously disappeared after approximately 3 weeks.

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Discussion

EP usually presents as an erythematous papules with a perilesional pale halo that disappears in a few days. This illness is unusual and the exanthem eruption is usually asymptomatic or slightly pruritic [2,8]. The first time that this disease was described was in 1969 by Cherry et al in children with enterovirus-enteric cytopathic human orphan (ECHO) virus infection [1] and Navarro, et al. [6] described a case report with clinical evidence of Epstein-Barr virus (EBV) in 1993, as well as, Prose et al described an eruption with similar features in 3 children, but unfortunately could not confirm its origin [2]. It is usually overserved in childhood [2,8], but in 2000, 10 cases were reported in Spanish adults, most of them in women [6,8]. In 2016, another case of a 73-year-old female was reported in Spain [9]. Recently, in 2018, 12 cases in adults were described from India by Chopra, et al. [10], who observed that females are more commonly affected. Our case report appears in summer, the same as the 7 cases described by Barrio, et al. [11]. The patient had no history of preceding symptoms. The onset of dermic lesions was 24 hours before consulting, which started from the extremities and spread to involve the trunk. The eruption was moderately pruritic. Multiple eruptive capillary hemangioma, viral exanthema, multiple pyogenic granuloma, popular urticaria or insect bites were considered in the differential diagnosis.

Histological findings show that epidermis is generally unaffected and there is no evidence of vascular proliferation; these findings justify the name of this disease: “eruptive pseudoangiomatosis” [2]. As we see in our patient, the clinical features and histological findings were similar to those described for EP. Its etiology is still unclear; however, most cases in the literature were related to viruses [6,12,13] but the development of the disease has also been observed due to insect bites5 and also in 2 subjects after taking an herbal medicine [13]. In our case report, we could perform a serology test and acute Parvovirus B19 infection was detected, which is first described in literature. Usually the family Paroviridae doesn’t affect humans, however, the only one that can be pathogenic in people is Parovirus B19. On the one hand, immunocompetent individuals’ immune system can control the infection, causing the erythema infectiosum (in children) and polyarthropathy (in adults). However, a pure red cell aplasia and chronic anemia can be caused in immunocompromised individuals [14]. Nevertheless, it has never been related to EP or to the onset of angioma-like lesions. There’s no specific treatment for EP, however, oral antihistamines and topical steroids can decrease symptoms [10]. As in our patient who received oral antihistamines.

Conclusion

This is the first case described in literature where the Parvovirus B19 is detected as the main cause of eruptive pseudoangiomatosis in adults. Because of that, it is necessary to keep in mind that many types of virus infection could cause this disease and we need to look for them conscientiously through laboratory and serological tests. Our report highlights the importance of keeping on studying the etiology of this disease in order to detect silent diseases, so that, we suggest that more cases need to be studied.

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Conflict of Interest

The authors have declared no conflict of interest.
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