Intertriginous bullous morphea: A clue for the pathogenesis?

Mukaddes Kavala, Ilkin Zindanci, Cuyan Demirkesen*, Emek Kocatürk Seyhan Beyhan, Zafer Turkoglu
Department of Dermatology, Goztepe Training and Research Hospital, Istanbul, *Department of Pathology, Istanbul University Cerrahpaşa Medical School, Istanbul, Turkey

Address for correspondence: Dr. Ilkin Zindanci, Goztepe Eğitim Hastanesi Dermatoloji Kliniği Kadıköy, İstanbul, Turkey.
E-mail: drilkinzindanci@yahoo.com.

ABSTRACT
Bullae occurring in lesions of morphea are uncommon. The cause of bullae formation in morphea is multifactorial, although lymphatic obstruction from the sclerodermatous process is considered the likeliest cause. Bullous morphea may be confused clinically with lichen sclerosus et atrophicus since both diseases may cause bullae in sclerodermatous plaques. A 69-year-old woman presented with a history of generalized morphea diagnosed 9 years earlier; and a 1-month history of pruritic bullae on her inframammary folds, axillary regions, lower abdomen, upper extremities and inguinal folds. Physical examination revealed multiple erythematous erosions, hemorrhagic vesicles and eroded bullae with slight scale or crusts overlying hypopigmented, indurated, shiny plaques. Skin biopsy revealed prominent edema in the papillary dermis, resulting in bulla formation and thickening of collagen fibers within the dermis. Direct immunofluorescence was negative. According to histologic and clinical features, the diagnosis of bullous morphea was established.

Key words: Bullous morphea, Intertriginous, Morphea, Pathogenesis

INTRODUCTION
Bullae occurring in lesions of morphea are uncommon, and the lower extremities are the most common sites of involvement.[1,2] The cause of bullae formation in morphea is multifactorial[3]; lymphatic obstruction from the sclerodermatous process is considered the likeliest cause.[3,4] Bullous morphea may be confused clinically with lichen sclerosus et atrophicus since both diseases may cause bullae in sclerodermatous plaques.[5,6]

CASE REPORT
A 69-year-old woman presented with a history of generalized morphea diagnosed 9 years earlier; and a 1-month history of pruritic bullae on her inframammary folds, axillary regions, lower abdomen, upper extremities and inguinal folds. The patient had no Raynaud’s phenomenon, dyspnea or dysphagia. Her current medication included moderate-strength topical steroid creams for the lesions. Physical examination revealed multiple erythematous erosions, hemorrhagic vesicles and eroded bullae with slight scale or crusts overlying hypopigmented, indurated, shiny plaques [Figures 1 and 2]. There were no facial telangiectasias or sclerodactyly. Blood count, erythrocyte sedimentation rate, creatinine, liver function tests and serum immunoglobulins were normal except 10% peripheral blood eosinophilia. ANA, anti-Ro, anti-La, Sm, dsDNA, Scl-70 and Borrelia antibody tests were negative. Serum protein electrophoresis increased in gamma-globulin fraction (23.5%; normal: 12-20%).

In skin biopsy, there was a prominent edema in the papillary dermis, resulting in bulla formation [Figure 3]. Collagen fibers within the dermis were thickened and had an eosinophilic, homogenous appearance. The skin appendages were entrapped within the coarse collagen bundles and were seen atrophic [Figure 4]. Direct immunofluorescence was negative. According to histologic and clinical features, the diagnosis of bullous morphea was established.

DISCUSSION
The existence of bullous lesions in morphea or systemic scleroderma is rare[1] and has been reported infrequently.[2,3,7,8]
Lichen sclerosus et atrophicus (LSA), in which the clinical features are confused with morphea, also forms bullae in the patches. The presence of bullae in LSA is more frequent than in morphea and has a hemorrhagic component. In the literature, there are conflicting reports about the nature and very existence of bullous morphea. Some authors have suggested that there is a close relationship between LSA and morphea, while others considered it as different manifestations of a single disease spectrum. However, Patterson and Ackerman emphasized that the two conditions were entirely separate.

Indeed, the histologic appearances of both diseases differ from each other. The epidermis in morphea shows neither hydropic degeneration of the basal cells nor follicular plugging, and the upper dermis in morphea has elastic fibers and shows no zone of edema. Reticular dermal changes of fibrosis and inflammation of morphea contrast with edema and loss of elastic tissue in LSA.

The etiology of bullous morphea is not certain, and there are several theories to explain the mechanism. Bullae formation was attributed to lymphedema caused by dilated lymphatic vessels, which occurs as a result of lymphatic obstruction from the sclerodermatous process. The tendency for bullae to form on the lower extremities also suggests that lymphatic obstruction, combined with increased hydrostatic pressure, leads to bullae. Pautrier also suggested that vascular changes like arteritis and phlebosclerosis play a role in bullae formation. Our patient had neither dilated lymphatic vessels nor vascular changes histologically. Su and Grene described a patient with bullous morphea profunda whose biopsy showed no lymphatic obstruction, arteritis or phlebitis. O’Leary found a correlation between local trauma and bullae formation. Dauod et al. showed the eosinophil granule component major basic protein (MBP) in the base of morphea bullae and suggested that eosinophils and, particularly, MBP may contribute to tissue destruction and bullae formation in some patients. Previous papers
have reported increased peripheral blood eosinophils and tissue eosinophilia in patients with morphea profunda and bullous morphea.2,20 Our case had significant peripheral eosinophilia but had no eosinophils in cutaneous tissue. The predilection of the intertriginous regions suggested the role of local trauma and friction as the precipitating factors of bullae formation.

Treatment of bullous morphea is unsatisfactory. It is suggested that salazopyrin has some efficacy.21 Daoud et al.2 found variable success with topical triamcinolone and hydroxychloroquine, 200 mg daily, but no improvement despite treatment with D-penicillamine, prednisone, cyclosporine, dapsone, minocycline, topical steroids, penicillin, cyclophosphamide and phenytoin. Since our patient refused systemic therapy, treatment was begun using a potent topical steroid cream. After 6 weeks, the blisters improved while the sclerotic patches persisted.

We conclude that as O'Leary had suggested before, the exact nature of blister formation on morphea lesions is due to friction, which becomes evident in the intertriginous regions of the body.

REFERENCES

15. Patterson JA, Ackerman AB. Lichen sclerosus et atrophicus is not related to morphea. A clinical and histologic study of 24 patients in whom both conditions were reputed to be present simultaneously. Am J Dermatopathol 1984;6:323-35.