Scleromalacia perforans secondary to a pemphigus.

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Abstract
Scleromalacia perforans is a rare form of anterior scleritis represented by progressive thinning of the sclera. It is most often associated with rheumatoid arthritis, and sometimes seen in inflammatory systemic disease. Rarely scleromalacia has been described in porphyria and herpes zoster. We are reporting the case of a 62-year-old man who was followed for Pemphigus vulgaris and who developed Scleromalacia perforans of the eye. The association between pemphigus and SP has never been described before. So this report suggests adding SP to rare but possible complications of pemphigus and highlights the importance of multi-disciplinary care.

Keywords: Scleromalacia perforans, Pemphigus, Ocular complications.

Introduction
Scleromalacia perforans (SP) is a rare form of anterior scleritis which readily presents as a blackish blue hue visible through a thin sclera [1]. No significant redness or pain is present but it is represented by progressive thinning of the sclera; It is a rare form of necrotizing anterior scleritis. It is often associated with systemic disease and involvement of multiple organs. We are reporting the case of a 62-year-old man who was followed for pemphigus vulgaris and who developed scleromalacia perforans of the eye.

This case is original because it is the first report of pemphigus patient who developed an SP.

Case Report
A 62-year-old hypertensive patient was followed in the dermatology department for a pemphigus vulgaris for 5 years treated with oral corticosteroid therapy with good initial development. The patient was lost from sight and then presented himself in a relapse chart of his pemphigus. On admission, the patient had diffuse cutaneous involvement with bullous lesions and post-bullous erosions. Otherwise, he reported a notion of eye pain with a decrease in the visual acuity of the right eye that had been evolving for 6 months. The examination objectified indeed a blue-blackish lesion of the right eye (Figure 1).

An ophthalmological examination was requested, showing the

![Figure 1. Scleromalacia perforans of the right eye.](image)
Discussion

Pemphigus is an autoimmune disease affecting the skin and mucous membranes characterized by the presence of antibodies against intercellular substance, which leads to suprabasal acantholysis and therefore blisters formation.

Ocular involvement can rarely be seen. It may be unilateral or bilateral and usually presents as chronic and refractory blepharitis and conjunctivitis [2,3]. Conjunctival bullae are unusual, [4,5] and sometimes, erosions and blisters over the lid margin and mucous discharge may be observed [6,7].

Persistent inflammation and erosion of the eyelids may affect the anatomy of the lid margin and the quality of the tear film and cause severe dryness, leading to poor vision quality. Atypical features are also reported, including cicatricial changes on both bulbar and tarsal conjunctivae [8], cobblestone-like conjunctival papillae [9], and trichiasis resembling mucous membrane pemphigoid [10]. No case of Scleromalacia perforans has been reported. Scleromalacia perforans is a rare severe disorder of the globe. It is a necrotizing scleritis without inflammation. Thinning and atrophy of the episclera are progressive without inflammatory signs, with the development of localized areas of scleral infarction. It is characterized by insidious and slow progression until changes of color sclera appear which corresponds to the choroid that appears through transparency through thin conjunctiva.

SP usually occurs in rheumatoid arthritis and more rarely during some vasculitis, systemic lupus erythematosus, Behçet disease, Wegener’s granulomatosis, it is also described in Crohn disease, porphyria, ankylosing spondylitis, and herpes zoster.

Young and Watson suggest three main causes leading to scleral destruction: prolonged local vasoconstriction, activation of scleral fibrocytes with resorption of the extracellular matrix, and infiltration of the scleral stroma by inflammatory cells, which explains the frequent occurrence of SP during inflammatory diseases [11]. The main complications are cataracts, anterior uveitis, glaucoma, and visual loss. Our patient had decreased visual acuity (6/10).

There is no therapeutic consensus. Treatment relies on corticosteroids and immunosuppressive drugs for severe necrotizing scleritis [12]. Surgical treatment is indicated in necrotizing scleritis with perforation of the globe. Surgery of SP is also needed when the uvea is exposed to preserve the integrity of the globe [11,13].

It is also necessary to treat or control the underlying pathology that led to SP to prevent recurrences [14]. For our case, the treatment of pemphigus allowed a stabilization of the ocular lesions.

There are some reports describing the effectiveness of infliximab in the treatment of scleritis in case of failure of immunosuppression [15,16]. Otherwise, Lola E et al. reported the successful treatment of two cases of refractory necrotizing scleritis with adalimumab [17].

As it is the first report of pemphigus patient who developed an SP, this report suggests adding SP to rare but possible complications of pemphigus.

Conclusion

In conclusion, despite the rarity of ocular involvement during pemphigus, it must always be sought in order to provide adequate care early and avoid complications. SP should be added to possible manifestations of pemphigus vulgaris.

Finally, autoimmune bullous diseases, especially pemphigus vulgaris, are serious pathologies with various complications and often require multidisciplinary management with the collaboration of dermatologists, ophthalmologists, and internists to provide optimal patient care.

References


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