To the Editor:

A 74-year-old man with a history of colon cancer and no history of sexually transmitted diseases presented with tender, moist, vegetating, and verrucous plaques localized to the inguinal creases and behind the scrotum of 3 weeks’ duration (Figure 1). The patient recently had taken lisinopril prescribed by his primary care physician for a couple of years for hypertension before switching to losartan prior to the current presentation. He later noticed the groin eruptions. He also noticed white tongue plaques temporally associated with the groin plaques and a long history of recurrent oral ulcerations. Prior to being seen in our clinic, outside physicians cultured methicillin-sensitive *Staphylococcus aureus* from the groin plaques and treated him with oral clindamycin, cephalexin, and topical mupirocin without a clinical response.

Our differential diagnosis included condyloma acuminata, condyloma lata, and cutaneous pemphigus vegetans. Laboratory testing revealed a nonreactive rapid plasma reagin test and peripheral eosinophilia of 14.9% (reference range, 0%–6%). Biopsy of a left groin plaque revealed epidermal hyperplasia with spongiosis and an eosinophilic-rich infiltrate on hematoxylin and eosin staining (Figure 2A), and direct immunofluorescence revealed diffuse epidermal intercellular IgG deposits (Figure 2B). The patient’s clinical and histologic presentation was consistent with cutaneous pemphigus vegetans. Biopsy of an oral ulcer revealed denuded acantholytic mucositis with eosinophilic-rich submucosal infiltrate and fibrosis (Figure 3A). Direct immunofluorescence was positive for lace-like intercellular staining for IgG and C3 within the squamous epithelium (Figure 3B). Together the clinical and histologic findings were consistent with oral pemphigus vulgaris.

The patient initially was started on oral minocycline 100 mg twice daily and mometasone furoate cream 0.1% twice daily to affected groin areas. With these interventions, the groin plaques almost completely resolved after several months, leaving only...
residual hyperpigmentation (Figure 4). The oral pemphigus vulgaris initially was treated with dexamethasone 0.5 mg/5 mL solution 2 to 3 times daily, but the lesions were refractory to this approach and also did not improve after the losartan was discontinued for several months. As such, mycophenolate mofetil was started. He was titrated to the lowest effective dose and showed near-complete resolution with 500 mg 3 times daily.

Cutaneous pemphigus vegetans, a rare variant of pemphigus vulgaris, is characterized by vegetating plaques commonly localized to the skin folds, scalp, face, and mucous membranes. Involvement of the oral mucosa occurs in a majority of cases. Although our patient had oral ulcerations, he did not have characteristic cerebriform changes of the dorsal tongue or associated verrucous hyperkeratotic lesions involving the buccal mucosa, hard and soft palate, or vermilion border of the lips that typically are seen in pemphigus vegetans. Subsequent biopsy of the oral mucosa confirmed oral pemphigus vulgaris in our patient.

This case presentation of co-occurring cutaneous pemphigus vegetans and oral pemphigus vulgaris is uncommonly reported in the literature. Although the etiology of this co-occurrence is not clear, it could represent a form of epitope spread, with the mechanism similar to that proposed for the progression of pemphigus vulgaris from the mucosal to the mucocutaneous stage by Chan et al, who suggested that an autoimmune reaction against specific desmoglein 3 epitopes (an important protein component for desmosomes and the autoantigen in pemphigus vulgaris) on mucosal membranes could induce local damage. These injuries could then expose the autoreactive immune cells to a secondary desmoglein 3 epitope present in the skin, leading to the development of cutaneous
lesions. Salato et al. also supported this idea of intramolecular epitope spread in pemphigus vulgaris, explaining that at various stages of the disease (mucosal and mucocutaneous), the antibodies have “different tissue-binding patterns and pathogenic activities, suggesting that they may recognize distinct epitopes.” This concept of epitope spread from the oral mucosal form to the cutaneous form of pemphigus vulgaris also could help explain our patient’s presentation, as he had a long history of recurrent oral ulcerations prior to developing the vegetating cutaneous plaques of cutaneous pemphigus vegetans.

We also appreciate that either the cutaneous pemphigus vegetans or oral pemphigus vulgaris could have been drug induced in our case. Captopril has been reported to cause pemphigus vulgaris, so it is conceivable that the related medication lisinopril was the culprit in our case. A prior case report described an elderly man who developed lisinopril-induced pemphigus foliaceus; however, there was no oral involvement in this case and no further blister formation within 48 hours of discontinuing lisinopril. An additional case report implicated lisinopril in the development of a bullous eruption on the oral mucosa in a female patient, though direct and indirect immunofluorescence did not reveal the autoantibodies that typically are seen in pemphigus vulgaris. Our patient’s blood eosinophilia also could support an adverse drug reaction.

Our patient’s losartan was discontinued for several months without respite of the oral ulcerations and thus was restarted. The cutaneous pemphigus vegetans continues to be in remission and was unaffected by restarting the losartan, making it a less likely culprit for his presentation.

We identified another case in the literature in which an individual with a history of colon cancer was diagnosed with cutaneous pemphigus vegetans. As such, we considered a possible link between the 2 diagnoses; however, the temporal disconnect between both conditions in our patient makes this less likely, unlike the other reported case in which the internal neoplasm and pemphigus vegetans appeared nearly simultaneously.

Finally, our case supports a combination of topical steroids and minocycline for treatment of cutaneous pemphigus vegetans.

Our case demonstrates the importance of considering cutaneous pemphigus vegetans in the differential diagnosis, despite its rarity, when patients present with vegetating plaques. In addition, although oral involvement is common with this condition, if the patient’s oral lesions do not fit the characteristic oral findings seen in pemphigus vegetans, alternative diagnoses should be considered.

REFERENCES