Molluscum Contagiosum Superimposed on Lymphangioma Circumscriptum

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To the Editor:
Lymphangioma circumscriptum (LC) is a benign malformation of the lymphatic system.1 It is postulated to arise from abnormal lymphatic cisterns, and it grows separately from the normal lymphatic system. These cisterns are connected to malformed dermal lymphatic channels, and the contraction of smooth muscles lining cisterns will cause dilatation of connected lymphatic channels in the papillary dermis due to back pressure,1,2 which causes a classic LC manifestation characterized by multiple translucent, sometimes red-brown, small vesicles grouped together. Lymphangioma circumscriptum can be difficult to differentiate from molluscum contagiosum (MC) due to the similar morphology.1 We present a notable case of MC superimposed on LC.

A 6-year-old girl presented with multiple grouped, clear, vesicular papules on the right buttock of 18 months’ duration. Some of the papules showed tiny whitish pearl-like particles on the top (Figure 1). Similar lesions were not present elsewhere on the body. She had no underlying disease and did not have a history of procedure, secondary infection of LC is common, with Staphylococcus aureus being the most common entity, but MC virus also can be secondarily infected.

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FIGURE 1. A, Multiple grouped, clear vesicular papules on the right buttock. Tiny whitish pearl-like particles were observed on top of the selected vesicular papules. B, Dermoscopy revealed whitish pearl-like structures and yellowish lacunae with minor vascular structures (original magnification ×10).
edema, or malformation of the lower extremities. Histopathology from one of the lesions showed dilated cystic lymphatic spaces in the papillary dermis lined with flattened endothelium and cup-shaped downward proliferation of the epidermis with presence of large intracytoplasmic inclusion bodies—features of both LC and MC (Figure 2). We waited 4 additional months for the MC lesions to self-resolve, but they persisted. The patient’s mother strongly requested for their removal, and the residual MC lesions were carefully removed by CO₂ laser. To prevent unnecessary physical damage to underlying LC lesions and minimize scarring, we opted to use the CO₂ laser and not simple curettage. She currently is under periodic observation with no signs of clinical recurrence of MC, but the LC lesions naturally persisted.

Due to its vesicular and sometimes warty appearance, LC can sometimes be hard to differentiate from MC. In one report, a vesicular plaquelike lesion on the trunk initially was misdiagnosed and treated as MC but was histologically confirmed as LC several years later.³ Our case demonstrates the coexistence of MC and LC. Although this phenomenon may be coincidental, we have not noticed any additional MC lesions on the body and MC only existed over the LC lesions, implying a possible pathophysiologic relationship. It is unlikely that MC might have preceded the development of LC. Although acquired LC exists, it has mostly been reported in the genital region of patients with conditions leading to lymphatic obstruction such as surgery, radiation therapy, malignancy, or serious infections.⁴ Because our patient developed lesions at an early age without any remarkable medical history, it is likely that she had congenital LC that was secondarily infected by the MC virus. Vesicular lesions in LC are known to rupture easily and may serve as a vulnerable entry site for pathogens. Subsequent secondary bacterial infections are common, with Staphylococcus aureus being the most prominent entity.¹ However, secondary viral infection rarely is reported. It is possible that the abnormally dilated lymphatic channels of LC that lack communication with the normal lymphatic system have contributed to an LC site-specific vulnerability to MC virus. Further studies and subsequent reports are required to confirm this hypothesis.

REFERENCES


FIGURE 2. Multiple dilated cystic lymphatic spaces in the papillary dermis lined with flattened endothelium. Cup-shaped downward proliferations of the epidermis with presence of large intracytoplasmic inclusion bodies also were observed (H&E, original magnification ×100).