Therefore, this case strongly supports that hypothesis. Rhode and Jennings\(^3\) described 4 types of Pacinian neuroma histologically, which are as follows: (1) a single enlarged Pacinian corpuscle, (2) a grape-like structure of normal-sized Pacinian corpuscles, (3) slightly enlarged Pacinian corpuscles arranged in tandem, and (4) hyperplastic Pacinian corpuscles arranged along the entire length of the digital nerve. Later, Reznik et al.\(^1\) considered types C and D as the same category. Important considerations within the differential diagnoses include neural-origin tumors such as schwannoma and glomus tumor. However, sometimes, other benign skin tumors such as mucoid cysts and fibromas could be considered as a differential diagnosis. The treatment of choice is surgical excision including the deep dermis and subcutis. In this case, the Pacinian neuromas appeared as exceptionally large, protruding masses, and we believe that they may have been induced by the accompanying dermal fibrosis and proliferation of the adnexal tissues. A literature review produced no reports of multiple and soft tumor-like Pacinian neuromas. Therefore, we herein report a case of uniquely presenting multiple Pacinian neuromas.

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An Unusual Presentation of a Progressive Zosteriform Macular Pigmented Lesion

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Dear Editor:

Rower et al.,\(^1\) defined progressive cribriform and zosteriform hyperpigmentation (PCZH) in 1978 as a disease that fulfills the following criteria: 1) cribriform pigmented macules that form a zosteriform distribution, 2) no history of skin disease or injury that would suggest postinflammatory hyperpigmentation, 3) an onset that arises well after birth, followed by gradual extension, 4) an onset that has no association with other skin diseases or internal abnormalities, and 5) characterized histologically by a mild increase in melanin pigment in the basal layer without nevus cells. In 1980, Simões and Piva\(^2\) described the progressive zos-
teriform macular pigmented lesion (PZMPL) as a chronic pigmentary dermatosis with similar clinical findings to PCZH. Although PZMPL is not completely understood, it is thought to be a variant of PCZH. PZMPL is distinguishable from PCZH because it presents with pruritus as a prodromal symptom and is characterized clinically by its abrupt spreading and histologically by pigmentary incontinence.

A 12-year-old boy presented with congenital brown pigmented skin lesions along Blaschko’s lines on the right side of his trunk and upper extremities (Fig. 1). He complained of occasional mild itching. The lesions had recently extended to the left side across the midline. The histopathological findings showed increased levels of melanin pigment in the basal layer and focal pigmentary incontinence in the upper dermis (Fig. 2).

The etiology of PZMPL is unknown. It has been speculated that its pathogenesis might be the same as other diseases that develop along Blaschko’s lines. Among the many inherited and some acquired skin conditions that develop along Blaschko’s lines, sporadic cases like the present one are often explained as inherited mosaicism. Further studies of similar cases are necessary to determine the pathogenesis of the disease.

PCZH arises after birth and follows the dermatomes. In terms of its hyperpigmentation and its asymptomatic nature, the case described in this report seems to align with the characteristics of PCZH, but the cribriform arrangement of small scattered macules is one particular characteristic of PCZH that differs from the extensive patches seen in the present case. The similarities of PZMPL to the present case include the presence of extensive plaques...
Giant Congenital Melanocytic Nevus with Proliferative Nodules Mimicking a Congenital Malignant Melanoma

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Dear Editor:

The malignant transformation of a giant congenital nevus (GCN) is extremely rare, and when it is discovered, the transformation may have already occurred during childhood. Benign proliferative nodules (PNs) in GCN may clinically and histologically mimic a malignant melanoma (MM), but clinically, PNs usually present at birth with multiple nodules, which is in contrast to MMs, and histologically, most patients with PNs do not present with progression of PZMPL. We recommend that dermatologists consider this disease when they encounter patients with chronic hyperpigmented dermatoses.

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