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Permalink
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Journal
Dermatology Online Journal, 22(12)

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Publication Date
2016

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Peer reviewed
Multiple pilomatrixcromas in the setting of myotonic dystrophy

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Abstract

The association between multiple pilomatrixcromas and the autosomal dominant neurodegenerative disorder myotonic dystrophy has been described in the literature. Although the mechanism is unknown, it is hypothesized that the dystrophia myotonica protein kinase mutation in myotonic dystrophy affects intracellular calcium levels, which alter proliferation and terminal differentiation that leads to cells that are observed in pilomatrixcromomas. We present a patient with multiple, symptomatic pilomatrixcromas and myotonic dystrophy, with a strong family history of both of these rare disorders.

Case Presentation

PATIENT: 30-year-old-man
DURATION: Over ten years
DISTRIBUTION: Left forearm and back

HISTORY: A 30-year-old-man presented to the Skin and Cancer Unit for evaluation of several, tender, firm, subcutaneous nodules that have been present for over ten years. The nodules were initially asymptomatic, but the lesions on his right upper back and left forearm have grown in size in the past three months and have become tender to palpation. The patient and his father do not recall ever having the lesions biopsied in the past, and the patient had never tried any treatments for the nodules.

On the day of presentation, one nodule on the left forearm underwent an excisional biopsy, and two weeks later, the nodule on his right upper back also was excised.

Medical history included myotonic dystrophy that was diagnosed at the age of 14 when he developed acute shoulder pain and weakness. His two sisters and his mother also have a diagnosis of myotonic dystrophy, and both of his sisters have multiple, similar skin lesions throughout their bodies as well. Other medical problems include obstructive sleep apnea. He does not have diabetes mellitus, cataracts, or heart failure although he has had a loop recorder implanted for the last 18 months to monitor for arrhythmias.

PHYSICAL EXAMINATION: Four subcutaneous, firm, mobile nodules were present on the left forearm and on the right upper, left upper, and left mid back. The largest nodule on the right upper back measured 4-cm x 2-cm with a linear hyperpigmentation that was consistent with a scar from a prior procedure.

Figure 1. Four subcutaneous, firm, mobile nodules were present on the left forearm and on the right upper, left upper, and left mid back. The largest nodule on the right upper back measured 4-cm x 2-cm with a linear hyperpigmentation that was consistent with a scar from a prior procedure.
left mid back. The largest nodule on the right upper back measured 4-cm x 2-cm with a linear hyperpigmentation that was consistent with a scar from a prior procedure (Figure 1). The other three nodules had no erythema, scale, or surface changes in the overlying skin. All of the lesions, in particular the largest nodule on the right upper back, were tender to palpation. The patient also has ptosis of the eyelids and dysarthria.

LABORATORY DATA: None

HISTOPATHOLOGY: There are aggregates of shadow cells in addition to nodules of basaloid cells. There is an admixed infiltrate of histiocytes, many of which are multinucleated (Figure 2).

DIAGNOSIS: Multiple pilomatricomas in the setting of myotonic dystrophy

Discussion
Pilomatricomas, which also are known as calcifying epitheliomas of Malherbe, are benign, cutaneous tumors that arise from the hair matrix and often occur as solitary, firm, mobile tumors on the head and neck of young women [1]. Multiple pilomatricomas in one patient are a rare finding. They have been associated with many disorders, such as Turner syndrome, Gardner syndrome, sarcoidosis, and the neuromuscular disorder myotonic dystrophy [2]. Myotonic dystrophy, which also is called Steinert Disease, is an autosomal-dominant disease with variable penetrance, and manifests as myotonia, lens opacities, mild mental retardation, frontal baldness, heart failure, and testicular atrophy. The association between myotonic dystrophy and multiple pilomatricomas was first described in 1965 [3]. Since then, there have been many reports of multiple pilomatricomas in several members of the same family, who also have myotonic dystrophy, which is similar to our patient [4-17].

The mechanism underlying the association between multiple pilomatricomas and myotonic dystrophy is unknown. Myotonic dystrophy is caused by a mutation in the dystrophin myotonica protein kinase (DMPK) gene that is located on chromosome 19. It is hypothesized that a mutation in the DMPK gene causes altered intracellular calcium levels, which play a role in cellular differentiation. In myotonic dystrophy, the mutated DMPK gene results in lower calcium levels, which causes high cell proliferation but lower terminal differentiation, and possibly leads to the shadow or ghost cells that are observed as a histopathologic feature in pilomatricomas [18]. Additionally, activating mutations in beta-catenin have been found in 75% of pilomatricomas, which may be implicated in myotonic dystrophy [19].

On histopathologic examination, pilomatricomas show sheets of epithelial cells of basophilic and eosinophilic types. Early pilomatricomas exhibit basophilic epithelial cells that cluster around the tumor periphery. As maturation occurs, these cells become more eosinophilic and lose their central nuclei and thus gain the appearance of characteristic shadow cells. Pilomatricomas also may exhibit central calcification that over time appears to resemble bony tissue [18]. The multiple pilomatricomas observed in myotonic dystrophy patients do not have distinct histopathologic features that vary from the solitary ones found in other patients.

Pilomatricomas rarely undergo malignant degeneration and are usually asymptomatic. Treatment typically is not warranted unless the lesions become symptomatic, often in the case
Case Presentation

of large pilomatricomas that cause pain due to ulceration or inflammation in the over-stretched overlying skin. Currently, the accepted treatment for pilomatricomas is complete surgical excision with clear margins, but for patients with several skin lesions, multiple excisions may leave many scars. The risk of recurrence after complete excision ranges from 0% to 6% [20]. Spontaneous regression has not been described.

Multiple pilomatricomas are a cutaneous finding in myotonic dystrophy, especially in patients with several family members, who exhibit both of these rare disorders. This association may be particularly helpful in patients who do not yet exhibit neurologic symptoms of myotonic dystrophy, as clinicians may then carry out the appropriate screening for this neurodegenerative condition.

References