Eccrine Hidrocystoma in a Child: An Atypical Presentation

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Eccrine hidrocystomas are small benign cystic lesions that normally affect only adult females. Eccrine hidrocystoma is characterized by tense vesicles that are predominately located on the face and particularly on the periorbital areas. Histologically, eccrine hidrocystoma consists of one or more partially collapsed unilocular cysts in the dermis, and these cysts are often situated adjacent to normal eccrine ducts. An 8-year-old boy presented with multiple skin-colored to bluish tiny papules on his nose. Histopathologically, the dilated partial cystic structures in the middermis were lined by one or two layers of flattened or cuboidal epithelial cells. We report here on an atypical presentation of multiple eccrine hidrocystomas that were localized on the nose of a child.

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INTRODUCTION

Robinson1 was the first to describe eccrine hidrocystoma (EH) in 1893. EHs characteristically appear as skin-colored to bluish vesiculopapular lesions, and most commonly on the periorbital area, and these lesions predominantly occur in middle-aged and elderly women2-3. EH is a benign small cystic lesion that usually enlarges during the summer and may disappear spontaneously in cooler weather2. To the best of our knowledge, EH in children has not previously been described in the medical literature and grouped EHs localized on the tip of the nose is a very rare and atypical presentation.

CASE REPORT

An 8-year-old boy presented with a 3-year history of multiple flesh-colored to bluish domed papules on the tip of the nose (Fig. 1). CO2 laser treatment had been performed at a local clinic during the prior year, but the lesions had not completely resolved. Both his past medical and family histories were not remarkable. The papules became more prominent in hot environments. The histopathology showed a dilated, partially cystic structure in the middermis (Fig. 2A). The cystic cavity was lined by one or two layers of cuboidal or flattened epithelial cells (Fig. 2B). No decapitation secretion was visualized. The immunohistochemical studies showed positivity for CEA, but negativity for S-100 and PAS (Fig. 3). The diagnosis of EH was established based on these
findings. Therapy with topical 1% atropine sulfate two times daily was instituted. The skin lesions showed great improvement after a month of treatment. No recurrence has been noted during the 1 year follow-up period.

**DISCUSSION**

EH usually presents as a small, translucent, cystic nodule that is 1 to 3 mm in diameter and the nodule often has a bluish hue. They are predominantly found on adult females, they occur as single or multiple lesions and they are mostly located on the periorbital and malar regions. EH is prevalent in adults between the ages of 20 and 70 years. There have been no reported cases of EH in childhood in the medical literature.

EH is currently classified into two types according to the number of lesions. The first type was described by Robinson and this is known as the Robinson type. Most of his patients were women who worked in hot and humid environments. Their lesions were characterized as having multiple small papules. Smith and Chernosky subsequently described a group of patients whose lesions were primarily solitary. The individual lesions of the multiple type are similar to the solitary lesions of the Smith type, except they are smaller. Tokura et al. suggested solitary EH is derived from the secretory coil, and the multiple ones are derived from the duct. The
Robinson-type multiple EHs are much more common in Korea and the Smith-type solitary EHs are relatively rare. The etiology of multiple EH remains unknown. A suggested cause for EH involves occlusion of the intradermal portion of the eccrine duct. However, Murayama et al. suggest that EH is a hamartoma-like disorder that becomes more pronounced as a result of retaining sweat. We thought that the latter hypothesis might explain the atypical, early presentation of our case.

EH must be differentiated from apocrine hidrocystomas. Apocrine hidrocystomas is usually larger and it does not become more prominent in hot environments or after exercise. Apocrine hidrocystomas affect the same age groups as do EHs, and they rarely occur during childhood or adolescence. Histopathologically, eccrine hidrocystoma, which is lined by ductal cells, differs from apocrine hidrocystoma by the absence of decapitation secretions, papillary projection into the lumen, PAS-positive granules and myoepithelial cells. EH is usually unilocular, whereas apocrine hidrocystomas are multilocular. Histochemically, solitary EH stains positive for S-100 protein, whereas the Robinson type EH and apocrine hidrocystomas do not. However, it may be difficult to differentiate EH from apocrine hidrocystomas even with conducting immunohistochemical study.

There is an interesting previously reported case of steatocystoma multiplex localized on the nose of a child. But the differentiation with EH was not difficult because steatocystoma usually has a yellowish color and it is histopathologically characterized by a folded cyst wall that consists of squamous epithelium, flattened sebaceous lobules within or close to the wall and a positive reaction of the lining cells for PAS. Clinically, other cystic lesions such as epidermal inclusion cyst, comedone and mucous cyst could be considered in the differential diagnosis, but all these lesions differ from EH histopathologically.

Although a solitary EH can be easily treated with surgical excision, eliminating multiple hidrocystomas is difficult because of their number and cosmetic location. The therapeutic options can be divided into medical and surgical modalities. Multiple lesions have been treated with topical scopolamine, topical atropine and botulinum toxin type A. Simple needle puncture has not been shown to produce lasting effects. Using a 585-nm pulsed dye laser has resulted in good outcomes when treating multiple EHs. Most importantly, avoiding hot temperatures or humid conditions will help prevent worsening of symptoms in patients. We chose topical atropine as a treatment method because of the patient's young age, and the number of skin lesions, and the site of presentation was cosmetically and functionally important.

The age at presentation and the clinical features of our case do not mimic the classic presentation of EH, be it the Robinson-type or the Smith-type. Due to the histologic findings, the lesion's aggravation during hot weather and the good response to atropine sulfate, we were able to make the diagnosis of EHs with an atypical presentation.

REFERENCES