Familial Eruptive Syringoma

Manal Elsayed¹, Magda Assaf²

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¹ Department of Dermatology & Venereology
² Department of Pathology Zagazig University, Egypt

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Abstract

Syringoma is a benign, eccrine sweat gland tumor. The typical clinical appearance of syringoma is firm, skin- colored to brown papules on the lower eye lids of middle - aged women. Familial syringoma is rare, but possibly is underestimated in prevalence. Eruptive syringoma, a separate entity, presents mostly in adolescents as clusters of numerous papules on the upper half of the body. Familial eruptive syringoma is a very rare clinical variant with only few reports in world literature.

Case study

Three sisters and two brothers presented to our Dermatology & Venereology department with disfiguring skin lesions asking for treatment. Their mean age was 27 years.

Case 1: A 23 year old female presented with asymptomatic lesions on abdomen, axillae, and neck (fig1)
Case 2 and Case 3: They are sisters of the first case (27, 36 year old respectively). They had similar distributed lesions on forearms, chest and axillae (fig 2, 3).
Case 4 and Case 5: They are brothers of the previous cases, of 22 and 32 year old respectively. The lesions were fewer than in female cases, on axillae, neck, chest and abdomen (fig 4).

On examination of their father, similar lesions appeared on lower eye lids, cheeks and anterior surface of neck. There is a history of consanguinity.
The lesions in all cases appeared as symmetrically distributed clusters of numerous skin colored and hyperpigmented rounded or flat-topped papules 1 to 3 mm in diameters. These lesions are asymptomatic first appeared at puberty, but additional lesions had developed later.

Biopsies were taken from different skin lesions in all patients. Results of histopathologic examination of lesions revealed aggregations of small tubular structures, lined by two rows of epithelial cells. Most of which are characterized by comma-like tails giving them a tadpole shape. They are embedded in a fibrous connective tissue stroma in the dermis (fig 5). These histopathologic findings were consistent with syringoma.

**Fig 5**: Photomicrograph showing numerous, scattered, dermal, small ducts lined by two rows of epithelial cells; some of which have epithelial cords with comma-like tails embedded in a sclerotic stroma (H&E x150).

**Discussion**

Syringoma is derived from Greek word syrnx, which means pipe or tube. It is a benign adnexal tumor of eccrine origin. It is seen commonly in females [1]. It is most common on the eye lids [2]. Other clinical variants, eruptive [3], linear [2], familial [4], vulvar [5], penile [6], scalp [3], acral [7] and plaque-type syringomas [8] have been described.

Eruptive syringoma is a clinical variant with many reports in world literature [3]. It is characterized by small, flesh-colored papules that occur in successive crops on the anterior body surface [9]. It arises in the peripubertal period. The lesions are bilateral, symmetrical, and have both follicular and non-follicular distribution [10]. In eruptive syringoma, a rare variant first described by Jacquet and Darier in 1987, the lesions occur in large numbers and in successive crops on the anterior chest, neck, upper abdomen, axillae, and periumbilical region at puberty or during childhood [6]. Eruptive syringomas are described more frequently
in patients with Down’s syndrome, and patients with Ehlers-Danlos syndrome [6]. In our cases no association have been detected.

Some literature has challenged the notion that eruptive syringoma is neoplastic. These authors maintain that eruptive syringoma actually represents a hyperplastic response of the eccrine duct to an inflammatory reaction. The pruritus which is sometimes associated with syringomas is another argument in favor of this hypothesis, while in our cases pruritus is absent. This hypothesis makes it difficult, however, to explain the familial occurrence of syringoma. Most authors still believe syringoma to be of sporadic or hereditary etiology [10,11,12]. Reed [13] described a family in which 7 females and 1 male in 4 generations were affected. Familial syringoma is rare, but possibly is underestimated in prevalence. In most reported families, syringoma have been observed to occur in an autosomal dominant pattern [14]. While familial eruptive syringoma is seldom reported, one author reported a family (mother, daughter and son) with dominantly inheritance eruptive syringoma. On contrary to our cases, syringomas were not presented on eye lids of any one of this family [15]. To the best of our knowledge, familial eruptive syringoma is very rare with only few cases have been reported.

In most reported families, syringomas have been observed to be more common in females. This female preponderance in both sporadic and familial forms of syringoma is well recognized but remains unexplained [4]. An association of syringoma with hormonal factors has been suggested; in particular the size of vulval syringomas may increase during pregnancy [6].

The lesions of syringoma may be disfiguring and often pose significant cosmetic concerns for patients [16]. There is no satisfactory treatment for wide spread syringoma. Surgical or chemical destruction involve some risks of scarring. Many treatment modalities such as dermabrasion, electrodessication with curettage and scissors excision have been tried with some success in limited sites [17]. Because they are located in the dermis and often numerous, unfortunately, different modalities of therapy yield poor cosmetic results. There is no standard treatment for eruptive syringomas. Although carbon dioxide lasers, topical tretinoin and trichloroacetic acid have been found to be useful, none eliminates the possibility of recurrence. Syringoma may also undergo spontaneous regression with age [3]. The patients presented here asked for treatment, but after discussion of known possibilities, they refrained from treatment.

References


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