On examination, discrete 2- to 3-mm perifollicular papules localized to the bilateral axillae (fig. 1) and the pubic area were noted. The color of the papules ranged from flesh-colored to slightly hypopigmented. Additionally, her left axilla showed areas of superficial erosion, erythema and overlying honey-colored crust (fig. 2). Of particular interest, the patient’s father was also examined in our office because of a similar eruption in both axillae, which was asymptomatic.

A diagnosis of Fox-Fordyce disease with superficial impetigo was made clinically, and the patient was treated with oral antibiotics and bacitracin ointment. The impetigo improved, but the apocrine miliaria persisted. Since the lesions were asymptomatic, she desired no further treatment.

**Discussion**

Fox and Fordyce first described the disease in 1902 [1]. They reported 2 cases – a young woman and a young man who presented with small papular lesions in the axillary region. The papules were numerous, firm, smooth, rounded and of normal color. Both patients experienced intense pruritus that affected their daily life. The histology of the lesions was described as hyperkeratosis involving chiefly the sweat duct orifices and their intraepidermal portions as well as a chronic dermal infiltrate.

In 1956, Shelley and Levy [2] further elaborated the description of Fox and Fordyce. They described it as occurring in the axillae, pubic area, labia, perineum, areolae, presternal area, umbilicus and the medial aspect of the upper thigh. Papules were
described as discrete, flesh colored and perifollicular with a smooth, dome-shaped contour, frequently with a central punctum. It is a disease most common in women between 13 and 35 years of age [2] and rarely seen before puberty [3], although more recent literature believes that the diagnosis is often missed in prepubertal girls [4]. As such, our case was initially misdiagnosed as genital warts.

The pathophysiology consists of a keratin plug in the terminal apocrine duct. This causes the dilatation of the apocrine sweat gland unit, and leads to ductal rupture and formation of an apocrine sweat retention cyst in the epidermis. Thus, an apocrine anhidrosis is noted. Pruritus is thought to be caused by extravasation of sweat [2]. The exact pathogenesis of the disease remains unknown although emotional factors, hormonal influences and chemical changes in the sweat components are all thought to be contributors [5].

Several treatments have been described with varying success. Most recently, pimecrolimus was shown to be successful in 3 young females [6]. Other methods mentioned in the literature include adapalene gel [4], liposuction-assisted curettage [7], topical clindamycin [8], oral retinoids [9], topical tretinoin cream [10, 11], topical 5% benzoyl peroxide combined with loratadine [12], ultraviolet radiation [13] and electrocoagulation [14].

Our patient’s father also exhibited classic Fox-Fordyce lesions. Rarely has Fox-Fordyce disease been mentioned as a familial disease, although 2 cases have been reported in sisters [8] and in 14-year-old identical male twins [15].

Conclusion

We present the 3rd case supporting a hereditary component in Fox-Fordyce disease and the first case involving two generations and opposite sexes. We suggest that a genetic component may be involved in the pathogenesis of the disease.

References