CASE LETTER

Papillary syringocystadenoma in an uncommon location

Dear Editor,

Papillary syringocystadenoma is a rare neoplasm of sweat glands, which is present at birth in 50% of cases.\(^1\) It has predominantly apocrine differentiation, although an eccrine origin has been described in some reports.\(^2,3\)

It usually presents as papule or plaque with a crustal surface that occurs almost exclusively in the head and cervical region.\(^4\)

This report details the clinical observations as well as the dermatoscopic and histopathological findings of a case in an unusual location.

A 6-year-old male, a native and resident of São José dos Campos, SP, presented a lesion with progressive growth five years ago on the left flank. At the dermatological examination, a papule of pink-erythematous coloration was observed, with a smooth surface and a fibroelastic consistency, measuring 5 mm × 3 mm (Fig. 1). Dermoscopic examination showed rounded structures of whitish-yellow color separated by whitish linear structures on an erythematous background (Fig. 2). The patient was referred for excision. Histopathological examination revealed the following: cystic invaginations covered by cells, sometimes squamous and sometimes columnar, with papilliferous projections to the light; tubular glands with large lights, covered by apocrine cells (Fig. 3).

Papillary syringocystadenoma (SCAP) is a rare adnexal tumor most often derived from apocrine cells.\(^3,4\) It predominates in children and adolescents, and is observed at birth in 50% of cases,\(^1\) which differs from the case reported above.

In 75% of cases, it is located on the head or cervical region. Some cases in other topographies have already been described (scrotal region, vulva, back, abdomen, and axilla). These locations, as well as that of the case described, are even rarer. When located on the scalp, it may be associated with the sebaceous nevus of Jadassohn.\(^4\)

Despite variable clinical presentation, a papule plaque is the most commonly found lesion type. In the majority, it is asymptomatic, but it can present pruritus, pain, and/or bleeding; it usually presents progressive growth, as in the case described.\(^1,4\)

To date, articles on the dermoscopic findings of SCAP are scarce. There is a description of a horseshoe vascular pattern, which was not observed in the present case.\(^1,5\)

SCAP can infect, bleed, ulcerate, and, in rare cases, it can progress to basal cell carcinoma (9%) or papillary syringo-cystadenocarcinoma. For these reasons, it was decided to perform the excision.\(^5\)

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\(^2,3\) Study conducted at the Dermatology Service, Santa Casa de São José dos Campos, São José dos Campos São Paulo, Brazil.
Conflicts of interest
None declared.

References
1. Karg E, Korom I, Varga E, Ban G, Turi S. Congenital syringocystadenoma papilliferum. Pediatr Dermatol. 2008;25:132–3.
2. Townsend TC, Bowen AR, Nobuhara KK. Syringocystadenoma papilliferum: an unusual cutaneous lesion in a pediatric patient. J Pediatr. 2004;145:131–3.
3. Bruno CB, Cordeiro FN, Soares FES, Takano GHS, Mendes LST. Aspectos dermatoscópicos do syringocistoadenoma papilífero associado a nevo sebáceo. An Bras Dermatol. 2011;86:1213–6.
4. Jalikh AP, Menezes AC, Gadelha A. Syringocistoadenoma papilífero localizado na coxa. Surg Cosmet Dermatol. 2013;5:273–5.
5. Einecke YS, Pinto EB, Silveira SO, Santos MA, Mendes AM, Carneiro FR. Syringocistoadenoma papilífero congênito. Rev SPDV. 2018;76:79–82.

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Multiple adult xanthogranuloma

Dear Editor,

Xanthogranuloma (XG) is a normolipemic non-Langerhans cell histiocytosis (NLCH) most commonly seen in childhood and generally designed as juvenile xanthogranuloma (JXG). Infrequently, XG can occur in adulthood. Both in adults and children, XG usually presents as a solitary lesion. Multiple lesions are rare in JXG and exceptional in adults. The authors report a case of multiple adult xanthogranuloma (MAXG) of the face, neck, trunk, abdomen, and axillae, with no extracutaneous involvement and no association with hematologic disease.

A 38-year-old female, with no relevant personal or family history of the disease, was seen in October 2016 with multiple papules on her skin, which she first noticed 18 months before. Dermatologic examination showed more than 100 yellowish-brown, smooth, firm papules with diameters of 1–3 mm on the face, neck, thorax, abdomen, and axillae (Fig. 1), two larger elements, standing out in the outer corner of the right eye and in the homolateral nasogenian groove. The lesions were asymptomatic and the general

Figure 1 Multiple xanthogranulomas. Detail of yellowish-brown, smooth, firm papules with diameters of 1–3 mm, asymptomatic, located on the face.