Triquilemoma Desmoplástico em Nevo Sebáceo do Couro Cabeludo

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RESUMO – Descrito originalmente por Jadassohn em 1895, foi apenas em 1932 que a designação “nevo sebáceo” foi introduzida, por Robinson. Trata-se de um hamartoma congênito relativamente prevalente que, classicamente, evolui por fases de crescimento e maturação (classicamente estão descritas 3 fases: infância <12; adolescência 12-18; adultos >18) e que, do 4º ao 7º decénio pode originar, em 10-30% dos casos, várias neoplasias epiteliais. Muito raramente reportados neste contexto, os triquilemomas são tumores anexiais benignos que se originam ou diferenciam no epitélio da bainha radicular externa folicular. Habitualmente solitários, podem apresentar-se clinicamente como pápulas translúcidas ou queratósicas faciais, em particular na proximidade da pirâmide nasal e dos lábios. A rara variante histológica “triquilemoma desmoplásico” pode simular clinicamente basaliomas (CBC), carcinomas espinhócélulares (CEC), nevos intradérmicos, verrugas virais, cornos cutâneos ou hiperplasias sebáceas e, histologicamente, evocar variantes desmoplásicas de CBCs, CECs e de carcinomas triquilêmicos. É apresentado caso de doente eurocaucasiano de 56 anos de idade que apresentava pápulas e nódulos translúcidos assintomáticos, desenvolvidos num nevo sebáceo do couro cabeludo. O estudo histológico revelou uma proliferação lobular com as características clássicas do triquilemoma desmoplásico no seio de um hamartoma sebáceo. O caso é ilustrado iconograficamente e a literatura relevante é sumariamente revista.

PALAVRAS-CHAVE – Nevo; Neoplasias das Glândulas Sebáceas; Neoplasias da Pele.

Desmoplastic Trichilemmoma arising on a Nevus Sebaceous of the Scalp

ABSTRACT – Originally described by Jadassohn in 1895, it was not until 1932 that the term “nevus sebaceous” was introduced by Robinson. It is a prevalent benign congenital hamartoma that classically evolves through discrete phases of growth and that, from the 4th to the 7th decade of life, can give rise in approximately 10 to 30% of cases to several epithelial neoplasms. Very seldom reported in this context, trichilemmomas are benign appendageal tumours that arise from or differentiate toward the outer root sheath epithelium of the normal hair follicle. Usually solitary, they present as verrucous, keratotic or smooth papules on the face, especially in the vicinity of the nose and lips. On clinical grounds, desmoplastic trichilemmomas, a rare benign histological sub-type of trichilemmomas, are easily taken for basal cell carcinomas, intradermal nevi, verrucae, cutaneous horns, sebaceous hyperplasia and squamous cell carcinomas; as far as pathology, differential diagnosis must include the desmoplastic variants of squamous and basal cell carcinomas, as well as trichilemmal carcinoma. A 56 year old Caucasian male presented with several asymptomatic pearly, translucent nodules and papules that had developed over the last 6 months within a congenital alopecic yellowish plaque of his scalp previously diagnosed, on clinical grounds, as a nevus sebaceous. Histopathological examination allowed to disclose a lobular neoplasm with the classic features of desmoplastic trichilemmoma. The case is iconographically depicted and the relevant literature is reviewed.

KEY-WORDS – Nevus; Sebaceous Gland Neoplasms; Skin Neoplasms.

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INTRODUCTION
Described by Headington and French in 19621 trichilemmomas (Tr) are benign tumours differentiating toward to the hair follicle isthmus. In 1990, Hunt at al2 described a desmoplastic histological variant characterized by a pseudocarcinomatous silhouette that often led to the misdiagnosis of desmoplastic squamous cell carcinoma or basal cell carcinoma. which evoked an invasive desmoplastic squamous cell carcinoma. Clinically, trichilemmomas may present either as solitary non-descript verrucose small papules at the central part of the face or, rarely, as multiple lesions at the same location, a circumstance where – together with sclerotic fibromas and acrokeratosis verruciformis - the diagnosis of Cowden’s syndrome should be evoked. Trichilemmomas may also, although rarely, occur as a secondary neoplasm within the milieu of a nevus sebaceous (NS)3 the very similar way varied appendageal tumours do, in what is regarded by some as a third stage in the natural history of NS.4,5. Pathologically, desmoplastic trichilemmoma (DTr) consists of well-defined lobules extending from the epidermis into the dermis centered by a dense, abundant hyaline stroma. In most cases, the epithelial lobules show at their margins the typical features of Tr lobules, with polygonal, PAS+ clear cells exhibiting peripheral nuclear palisading that lie upon a thickened eosinophilic hyaline membrane. As the center of the lesion is approached, the epithelial sheaths progressively get narrower, cells diminishing in dimension and becoming more and more irregular. When cells merge on the dense stroma, the lobular pattern is disrupted and cells tend to cluster in irregular cords or nests, with occasional dyskeratosis or necrosis, thus simulating an invasive carcinoma.2,3,6 The case of a 56 year old Caucasian male who developed a desmoplastic trichilemmoma within a nevus sebaceous of the scalp is presented.

CASE REPORT
A 56 year old Caucasian male presented with several asymptomatic papules that had developed over the last 6 months within an alopecic plaque on the scalp, previously diagnosed, on clinical grounds, as a sebaceous nevus. He denied any previous local major trauma, radiation therapy or topical agents as well as any family or personal history of skin or visceral cancer. He had lived his whole life in Angola but, due to his fair complexion, had always avoided deliberate or any kind of recreational sun exposure.

On physical examination, the patient had several papules – from 3 to 6mm diameter – superimposed, in an apparently multifocal distribution, on a 15X12 mm diameter, alopecic, slightly raised yellowish plaque with a smooth, velvety surface (Fig. 1). The papules were hemispheric or pedunculated, some warty and some translucid. The examination of the rest of the integument was nondescript.

The patient declined to undergo excision of the entire mass. A biopsy was then suggested for diagnostic purposes, which was readily accepted. A large shave biopsy that involved the whole lesion was then performed, with subsequent CO2 laser vaporization of the resulting open wound, for the sake of hemostasis.

Histological examination revealed an endoexophytic epithelial proliferation that filled the dermis and was contiguous to the overlying acanthotic epidermis (Fig. 2). The neoplasm was composed of well-demarcated lobules, with two populations of epithelial cells: one, centrally located, with clear and abundant cytoplasm (Fig. 3) and the other, peripherally, with basophilic cells with nuclear palisading lined by a thickened eosinophilic hyaline membrane (Fig. 4). Overall, a dense hypercellular and desmoplastic stroma occupied the central portion of the proliferation (Fig. 5). There was neither evidence of cytological atypia nor of mitotic figures. At the extreme periphery of the histological section, one could appreciate some trabecular acanthosis of the epidermis, hyperplastic sebaceous glands – some opening directly to the surface – and a focal lymphomononuclear inflammatory infiltrate (Fig. 6).

The diagnosis of desmoplastic trichilemoma arising in a sebaceous nevus was made. The tumour appeared to be fully confined within the limits of the tissue specimen, suggesting...
that a complete excision had been performed. Meanwhile, the resulting surgical wound, which measured 10x6mm diameter, fully healed by second intention, in a three-week period without major pain or complications. At follow-up, 12 months afterwards, there was no recurrence of the neoplasm (Fig. 7). Although fully informed about the possible development of other local neoplasms, the patient nevertheless refused to undergo complete excision of the congenital sebaceous nevus.

**DISCUSSION**

Originally described by Jadassohn in 1895, it was not until 1932 that the term “nevus sebaceous of Jadassohn” (NS) was introduced, by.  

It refers to a well-recognized congenital hamartoma of the skin that appears as a slightly raised, yellowish, smooth alopecic plaque on the scalp, face and neck. It evolves through distinct phases of growth, clearly accelerated at puberty, when it may show a hyperplastic, nodular appearance and, at adulthood (4th-7th decades), where can give rise,
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in approximately 10-30% of cases, to diverse epithelial neoplasms. Several reported activating mutations in HRAS and KRAS have been reported in nevus sebaceous, so as to allow to encompass by some authors it as the result of a proliferative whole skin somatic mosaic status. Most of these tumours are benign and have a pilar, sebaceous, eccrine or apocrine origin. Malignant neoplasms are extremely rare among children as well as in adults, as it has been shown that most of the lesions formerly classified in this setting as basal cell carcinomas (BCC) were actually trichoblastomas.

Since its original description in 1990, desmoplastic trichilemmomas (DTr) have very seldom been reported, particularly in the setting of NS. The total of only 81 cases reported worldwide up to 2012 may in fact represent an underestimation of the actual prevalence of these tumours. It has been reported to occur in several ethnicities, seeming to be more prevalent in males and to favor the 5th-6th decades of life. As far as location, DTr’s seem to predominate on the facial area, although extrafacial locations have also been described. Clinically, its presentation is quite non-specific, either as a lobular or diffuse thickening of a previous NS or as newly superimposed translucid or hyaline connective tissue. The amount of desmoplasia is said to vary, but in most cases represents about 20-60% of the tumour. It is important not to misdiagnose this pseudoinvasive pattern as an invasive carcinoma. Although some cytological pleomorphism, individual dyskeratosis and cellular necrosis may focally be present, overall, the absence of frank cell atypia and mitotic activity do favor a diagnosis of DTr. In difficult cases however, the CD 34 positive and BerEP4 negative immunostaining of the epithelial cells allow to definitely clarify diagnosis and rule out BCC.

In our patient’s case, as opposed to the non-specific clinical presentation, the pathological examination allowed to clearly diagnose DTr within a nevus sebaceous. As far as therapeutic intervention in NS, a lot of controversy does exist in the literature. Prophylactic surgery during childhood has classically been advocated at the light of the natural history of NS and the early assumed risk of malignancy, namely basal cell carcinoma, as the lesion ages. As opposed to that view, others have suggested that prophylactic surgery should not systematically be performed due to the more modern evidence of its low malignancy risk. In the context of these pros and cons however, we do believe that the decision should be the result of a careful individual based, case-by-case consideration, whereby several facts should be weighed: 1. Secondary neoplasms (mostly benign) are a common occurrence in NS’s: 2. On clinical grounds, changes in long lasting NS’s of the scalp are hardly if ever noticeable; 3. Secondary neoplasms often lack clinical specific features as to allow to clinically differentiate between benign or malignant; 4. Some molecular evidence does exist in that NSs share with BCCs a defect of the tumour suppressor gene PTCH (9q22.3), which might be in keeping with a definite carcinogenic potential of NS (20). Thus, having these data in mind, we suggest that at least in most of the NS’s of the scalp area, surgical excision is indeed advisable. That was not unfortunately our patient’s case, in whom due to personal schedule limitations, we were only able to perform a large (excisional) biopsy of the desmoplastic trichilemmoma, without being able to completely remove the nevus sebaceous.

In what concerns DT treatment, although usually benign, its potential coexistence with atypical basaloid cell proliferations including CBC, makes Mohs surgery (for both adequate margin control and tissue sparing benefits) or complete surgical excision with pathological confirmation of clear margins the treatments of choice.

In summary, a case of a patient with a desmoplastic trichilemmoma arising in a nevus sebaceous of the scalp is reported. Diagnostic and therapeutic challenges posed by this rare tumour when in the context of a complex, still enigmatic skin hamartoma such as nevus sebaceous are discussed.

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