

Desmoplastic trichoepithelioma on the face mimicking basal cell carcinoma: a case report and literature review

Tricoepitelioma desmoplásico na face simulando carcinoma basocelular: um relato de caso e revisão da literatura

Isabele M. Saldanha^{1*}, Airtón K. Motizuki², and Renata M. Oyama-Okajima¹

¹Dermatology Service of the Federal University of Pará, João de Barros Barreto University Hospital; ²Department of Medicine, Faculty of Medicine of Federal University of Pará. Belém, Pará, Brazil

Abstract

Desmoplastic trichoepithelioma is a rare benign cutaneous neoplasm that differentiates toward germinative cells of the hair follicle. It can be classified as familial multiple trichoepithelioma, solitary, and desmoplastic, the latter being a rare variant with low incidence and clinical features resembling basal cell carcinoma. We report the case of a 54-year-old female patient who presented with a brownish plaque exhibiting areas of dark pigmentation, an irregular outline, a pearly sheen, and surface telangiectasias in the right paranasal region. Complementary studies, including histopathological and immunohistochemical analyses, were performed to establish the definitive diagnosis. Furthermore, this cutaneous lesion displays both clinical and histopathological characteristics that complicate differentiation between benign and malignant tumors, making immunohistochemistry essential to confirm the desmoplastic variant of trichoepithelioma.

Keywords: Desmoplastic trichoepithelioma. Differential diagnosis. Basal cell carcinoma.

Resumo

O tricoepitelioma desmoplásico é uma rara neoplasia benigna de pele que se diferencia em células germinativas do folículo piloso, e pode ser classificado entre tricoepitelioma múltiplo familiar, solitário e desmoplásico, sendo esse último uma variante rara com pouca incidência na população e com similaridades clínicas do carcinoma basocelular. O caso é de uma paciente de 54 anos, que apresentou uma placa acastanhada, com algumas áreas de pigmentos enegrecidos, irregular, com brilho perláceo e telangiectasias em sua superfície na região paranasal direita, sendo realizado exames complementares como o anatopatológico e imunohistoquímico para confirmação completa do diagnóstico. Além disso, essa lesão no tegumento manifesta estruturas tanto clínicas quanto histopatológicas que dificultam na diferenciação de tumores malignos ou benignos, sendo fundamental a realização da imunohistoquímica para confirmar essa variante desmoplásica do tricoepitelioma.

Palavras-chave: Tricoepitelioma desmoplásico. Diagnóstico diferencial. Carcinoma basocelular.

*Correspondence:

Isabele M. Saldanha

E-mail: isabelemartinsaldanha@gmail.com

2795-501X / © 2025 Portuguese Society of Dermatology and Venereology. Published by Permanyer. This is an open access article under the CC BY-NC-ND license (<http://creativecommons.org/licenses/by-nc-nd/4.0/>).

Received: 28-07-2025

Accepted: 05-10-2025

DOI: 10.24875/PJDV.25000049

Available online: 28-10-2025

Port J Dermatol and Venereol. 2026;84(1):46-49

www.portuguesejournalofdermatology.com

Introduction

Desmoplastic trichoepithelioma (DT) is a rare benign skin neoplasm of the hair follicle, a variant of trichoblastoma originating from the proliferation of mesenchymal cells¹⁻³. The classification of this cutaneous neoplasm can be divided into three forms: familial multiple trichoepithelioma, solitary, and desmoplastic, with the desmoplastic form characterized by increased fibroplasia in the dermis and this tumor should be considered a differential diagnosis for basal cell carcinoma (BCC)¹⁻³.

In addition, DT shows a predilection for females, particularly young or middle-aged women^{4,5}. Its most common manifestations occur on the facial skin as pink or skin-colored papules and nodules that progressively increase in size, often easily confused with the clinical aspects of BCC^{4,5}. The histopathology of this neoplasm consists of a histological triad of thin cords of basaloid cells, keratin cysts, and desmoplastic stroma, which are important for both histological differential diagnoses and clinical differentiation from BCC³⁻⁷.

Case report

A 54-year-old female patient was referred for a dermatological consultation due to a facial lesion that had been present for approximately 2 years, showing progressive growth over time without associated symptoms (Fig. 1).

On dermatological examination, a brownish plaque with some black-pigmented areas, irregular shape, pearly shine, and telangiectasias on its surface was observed, measuring 1 cm in the right paranasal region (Fig. 2).

The patient underwent an incisional biopsy of the lesion, followed by histopathological examination, which showed atrophic epidermis and dermis with diffuse proliferation of small nodular masses and cords of basaloid cells with peripheral palisading, including occasional horn cysts, embedded in desmoplastic and myxoid stroma. Bulb-papilla images suggested follicular differentiation (Fig. 3).

The histopathological examination revealed an intra-dermal epithelial neoplasm, raising differential diagnoses of DT, micronodular BCC, and microcystic adnexal neoplasm. An additional immunohistochemical study confirmed the diagnosis of DT (Table 1).

Following the DT diagnosis, clinical follow-up of the patient was chosen.

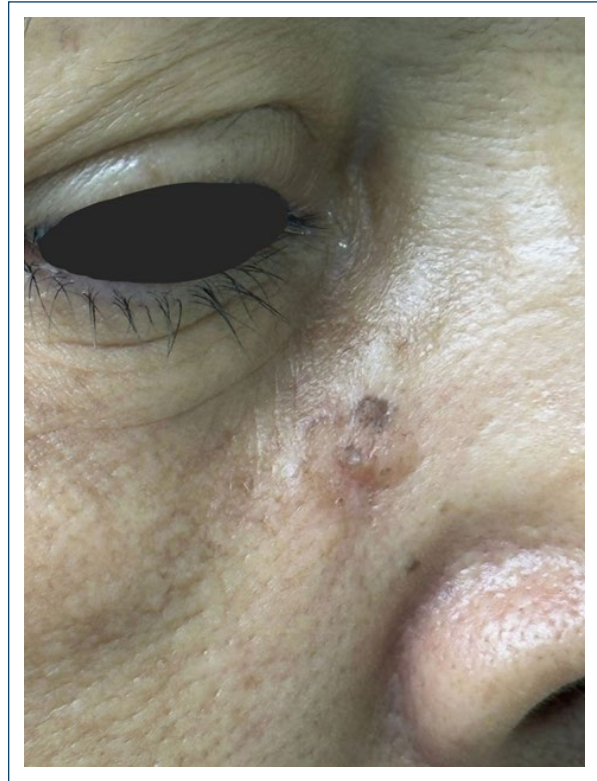


Figure 1. Clinical appearance of the neoplasm.



Figure 2. Dermatoscopy of the lesion with irregularly distributed black pigment, tortuous vessels, and mild erythema, measuring 1 cm.

Discussion

BCC is the most common skin neoplasm among non-melanoma skin cancers. Histopathological classification divides BCC into types such as nodular, superficial, and infiltrative. Despite its rare metastatic potential, BCC significantly impacts patient morbidity¹⁻³. DT is a

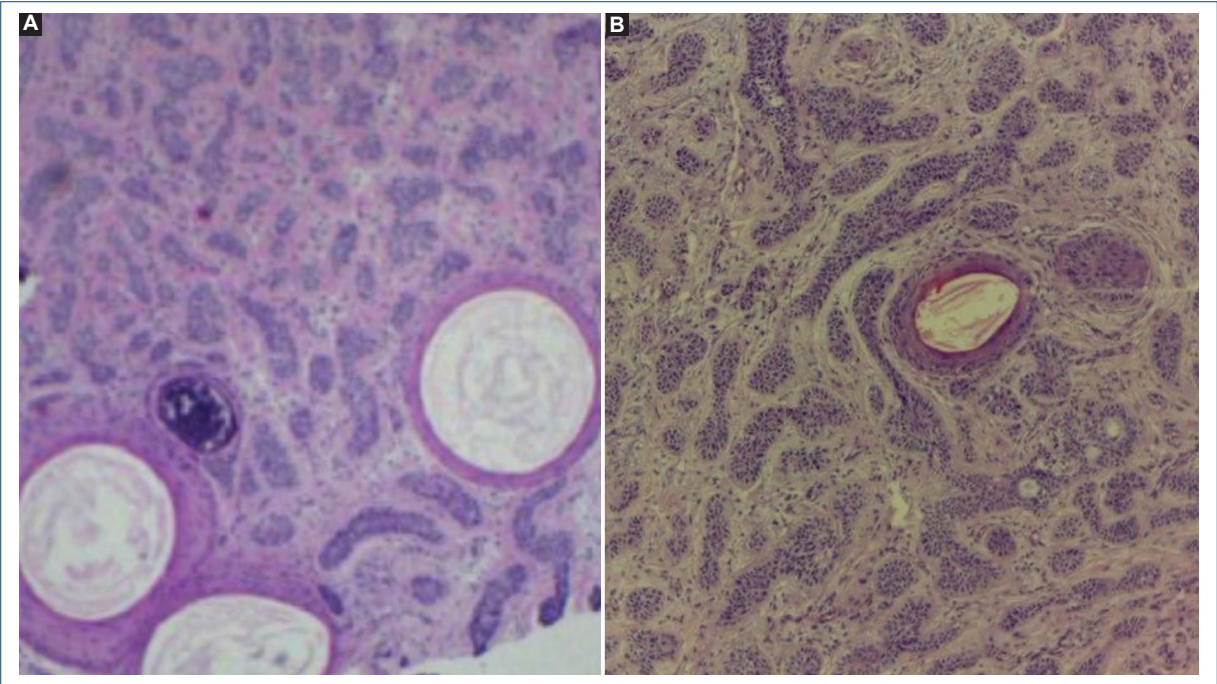


Figure 3. Histopathological findings. **A** and **B**: histological sections, horn cysts, embedded in desmoplastic and myxoid stroma (H&E, x40).

Table 1. Results of the immunohistochemical panel

Antibody	Clone	Result	Observation
Ki-67 (cell proliferation antigen)	MIB1	Positive	5%
p63 protein (squamous/transitional epithelia; myoepithelial cells)	DAK-p63	Positive	-
Epithelial tumor glycoprotein	BerEp4	Positive	-
Epithelial membrane antigen	E29	Negative	-
Anti-apoptotic protein BCL-2	124	Negative	-

benign cutaneous tumor that differentiates into germinative hair follicle cells¹⁻³. Representing < 1% of all skin tumors, its clinical and histopathological similarities with BCC components pose a controversy in both dermatologic surgery and clinical settings, as this neoplasm has a rare chance of malignant transformation³⁻⁷.

According to Rezze et al. dermoscopy is an essential technique in the clinical approach to skin lesions, playing a key role in identifying malignant lesions such as BCC⁸. However, dermoscopic findings in DT can mimic BCC structures, such as pearly shine and branching

telangiectasias⁹. While arborizing telangiectasias are common in many cutaneous neoplasms, these vessels are thicker and more branched in nodular BCC, unlike the finer, less branched vessels in DT, which aligns with this patient's dermoscopic findings⁹.

Histopathological findings in trichoepithelioma are characterized by basaloid cell proliferation, compact desmoplastic stroma, small cysts lined with keratinized epithelium, and sometimes calcifications. These features overlap with BCC, necessitating immunohistochemical studies for definitive diagnosis, as trichoepithelioma does not express carcinoembryonic antigen¹⁰⁻¹⁴.

Studies like Bains et al. used markers CK20, Ki-67, Bcl-2, CD10, and CD34 in immunohistochemistry to evaluate reactivity between the two neoplasms discussed. However, CD34, CD10, and CK20 markers were not used in this case; only Bcl-2 and Ki-67 were applied¹⁵. The Ki-67 marker showed 5-10% tumor cell proliferation, consistent with Bains et al.'s findings, while Bcl-2 did not show reactivity, highlighting the need to define an immunohistochemical panel to differentiate this benign tumor from other malignant neoplasms¹⁵.

Given this, there is limited scientific evidence on trichoepithelioma and its variants, especially the desmoplastic form, as it is a rare neoplasm with low population incidence. Only one publication exists in Brazilian literature.

Furthermore, few studies address specific markers for identifying and differentiating this disease. Thus, continuous monitoring of suspected DT lesions is crucial to correctly rule out malignant cutaneous diagnoses and implement appropriate therapeutic management.

Conclusion

The patient in this case report presents with DT, a rare but benign tumor that clinically and histopathologically resembles BCC. Immunohistochemistry is an essential and necessary test for diagnostic confirmation and for defining the appropriate therapeutic management. In addition, there is limited literature evaluating the desmoplastic form of trichoepithelioma and its histological or immunohistochemical diagnosis. Therefore, careful attention is needed to avoid diagnostic errors and unnecessary treatments for patients diagnosed with this condition.

Funding

None.

Conflicts of interest

None.

Ethical considerations

Protection of human subjects and animals. The authors declare that no experiments on humans or animals were performed for this research.

Confidentiality, informed consent, and ethical approval. The authors have followed their institution's

confidentiality protocols, obtained informed consent from all patients, and secured approval from the Ethics Committee. SAGER guidelines have been followed as applicable to the nature of the study.

Declaration on the use of artificial intelligence (AI). The authors declare that no generative artificial intelligence was used in the writing or creation of the content of this manuscript.

References

1. Mohammadi AA, Seyed Jafari SM. Trichoepithelioma: a rare but crucial dermatologic issue. *World J Plast Surg.* 2014;3:142-5.
2. Azulay, RD. *Dermatologia.* 8th ed. Rio De Janeiro: Guanabara Koogan; 2022.
3. Fitzpatrick TB. *Tratado De Dermatologia.* 7th ed. Rio De Janeiro: Revinter; 2010.
4. Rahman J, Tahir M, Arekemase H, Murtazaliev S, Sonawane S. Desmoplastic trichoepithelioma: histopathologic and immunohistochemical criteria for differentiation of a rare benign hair follicle tumor from other cutaneous adnexal tumors. *Cureus.* 2020;12:e9703.
5. Jedrych J, Leffell D, McNiff JM. Desmoplastic trichoepithelioma with perineural involvement: a series of seven cases. *J Cutan Pathol.* 2012;39:317-23.
6. Reiter O, Mimouni I, Dusza S, Halpern AC, Leshem YA, Marghoob AA. Dermoscopic features of basal cell carcinoma and its subtypes: a systematic review. *J Am Acad Dermatol.* 2021;85:653-64.
7. Simpson W, Garner A, Collin JR. Benign hair-follicle derived tumours in the differential diagnosis of basal-cell carcinoma of the eyelids: a clinico-pathological comparison. *Br J Ophthalmol.* 1989;73:347-53.
8. Rezze GG, Sá BC, Neves RI. Dermatoscopia: o método de análise de padrões. *An Bras Dermatol.* 2006;81:261-8.
9. Pitarch G, Botella-Estrada R. Dermoscopic findings in trichoblastoma. *Actas DermoSifiliogr.* 2015;106:e45-8.
10. Almeida HL, Boeing M, Parodi RB, Mendonça R. Tricoepitelioma desmoplásico. *An Bras Dermatol.* 2001;76:4.
11. Heller J, Roche N, Hameed M. Trichoepithelioma of the vulva: report of a case and review of the literature. *J Low Genit Tract Dis.* 2009;13:186-7.
12. Khan H, Mpyisi L, Azzam MJ, Braudis K. Infiltrative basal cell carcinoma (iBCC) adjacent to a desmoplastic trichoepithelioma (DTE): a case report and literature review of iBCCs and DTEs. *Cureus.* 2024;16:61893.
13. Brownstein MH, Shapiro L. Desmoplastic trichoepithelioma. *Cancer.* 1977;40:2979-86.
14. Huet P, Barnéon G, Cribier B. Trichoépithéliome desmoplastique: corrélation dermatopathologie-dermatoscopie. *Ann Dermatol Venerol.* 2019;146:760-3.
15. Bains A, Vishwajeet V, Lahoria U, Bhardwaj A. Morpheaform basal cell carcinoma masquerading as desmoplastic trichoepithelioma in a young patient. *Indian J Dermatol.* 2021;66:706.