Neurofolicular hamartoma presenting as a rare adnexal neoplasm in association with basal cell carcinoma

Abstract

Background: Neurofolicular hamartoma (NFH) is characterized histopathologically by fascicles of spindle cells that laterally delimited by hyperplastic folliculosebaceous units. It usually appears on face, near the nose or nasolabial fold. It does not manifest true neural differentiation and recently the term spindle cell predominant trichodiscoma (SCPT) has been used instead.

Case Presentation: We present a case of a 40-year-old male with co-incidence of NFH and basal cell carcinoma (BCC) that the mesenchymal components of NFH were similar to SCPT but these components highly expressed S100 protein. We also discuss about the histological aspect of the neoplasia in this report and consider the findings of other reports in association with classification of NFH by means of cellular markers and morphological resemblance to other skin hamartomas.

Conclusion: Neurofolicular hamartoma is a rare benign tumor that thought to represent the cellular end of a morphological spectrum with trichodiscoma. The morphological features and expression of S100 protein in neural element helped us to achieve the diagnosis of neurofolicular hamartoma. However, variable reports of S-100 protein expression in NFH are available and further studies are needed to determine the classification of this tumor.

Keywords: Skin tumor, neurofolicular hamartoma, Basal cell carcinoma, S-100 protein, immunohistochemistry

Citation:
Montazer F, Sanei Motlagh AR. Neurofolicular hamartoma presenting as a rare adnexal neoplasm in association with basal cell carcinoma. Caspian J Intern Med 2019; 10(1): 107-110.
Case Presentation

A 40-year-old male presented with the appearance of skin lesions on his face during the past two months. Past medical history of diabetes mellitus was noted for him without any genetic disorder. On clinical examination, two skin-colored dome shaped firm masses were noted with each size of 10×5mm and 5×5mm, respectively. The larger lesion showed surface ulcerations. With suspicion of basal cell carcinoma (BCC) for both lesions, excisional biopsy was performed and sent for pathological study including hematoxylin and eosin (H&E) and immunohistochemical (IHC) staining for S-100, neuron specific enolase (NSE), smooth muscle actin (SMA), and desmin markers. Histopathological results confirmed the BCC, nodular type, for the larger one (figure 1).

Figure 1. Hematoxylin and eosin (H&E) staining of the larger lesion showing basal cell carcinoma (BCC), nodular type

H&E stained sections of smaller lesion showed epidermis with unremarkable changes. The dermis displayed a well circumscribed, non-encapsulated lesion composed of epithelial and mesenchymal components. The epithelial component consist of distorted and hyperplastic pilosebaceous units with prominent sebaceous glands (fig 2).

Figure 2. Distorted and hyperplastic pilosebaceous units with prominent sebaceous glands. H&E staining of epithelial and mesenchymal components

The mesenchymal component mainly showed myxoid and fibrillary appearance containing elongated and wavy spindle cells arranged in fascicles resembling neurofibroma (figure 3).

Figure 3. Myxoid and fibrillary appearance containing elongated and wavy spindle cells (H&E staining)

Further immunohistochemical study for confirmation of neural mesenchymal stroma was done. Fibrillary mesenchymal components express S-100 marker (figure 4), while neuron specific enolase, smooth muscle actin, and desmin were negative (figure 5). Based on the results of H&E staining and IHC, the diagnosis of neurofollicular hamartoma was confirmed. Neurofollicular hamartoma itself is a benign tumor and treatment was achieved by local excision. The BCC lesion was demarcated and small in size which was surgically removed without further topical treatment or radiotherapy.

Figure 4. Immunohistochemical staining, fibrillary mesenchymal components express S-100 marker
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Figure 5. NSE and SMA, and desmin markers were negative in IHC staining of mesenchymal components

Discussion

Neurofolicullar hamartoma as a rare benign tumor is thought to represent the cellular end of a morphological spectrum with trichodiscoma. The clinical differential diagnosis includes fibrous papule, basal cell carcinoma, and dermal nevus (3). NFH is the tumor with follicular differentiation that expresses diffuse CD34 stromal positivity, Bcl-2 on outermost basal cells, and CD10 on perifollicular and peritumoral stroma. It shares these features with basaloid follicular hamartomas (BFH) and vellus hair hamartoma (VHH) which have the same follicular growth pattern (4). A case report of neurofolicullar hamartoma showed that these lesions could express strong and diffuse pattern of S100 protein on spindled cells. Scattered positivity of spindle cells for monoclonal neuron specific enolase and synaptophysin was also noted (5). The pathologic study results in our patient demonstrated that the lesion had myxoid and fibrillary components with wavy nuclei with focal S-100 positivity without any sign of neural differentiation. They used the term spindle cell predominant trichodiscoma (SCPT) instead of neurofolicullar hamartoma to describe precise morphological entity. Differential diagnosis of NFH should include all fairly circumscribed superficial cellular lesions composed of CD34 positive spindle cells. Despite clinicopathological resemblance of angiofibroma to NHF, angiofibroma shows a dense collagenous stroma with a diffuse scatter of single cells in conjunction with more capillary vessels (6). Kacerovska et al. reported a case of SCPT with a focal palisaded arrangement of stromal cell just like those seen in schwannoma, in which the stromal cells were positive for CD34 and negative for S-100 protein. The similarity of SCPT to peripheral nerve neoplasms was mentioned by illustrations (7).

Basal cell carcinoma (BCC) is the most common cutaneous malignant neoplasm. It has slow growth pattern and rarely metastasize but a small proportion (less than 0.5%) of this tumor is locally aggressive. Nests of uniform basaloid cells within the dermis that are often separated from the adjacent stroma by the thin clefts are apparent in H&E stained sections (8). The lesions in this patient were small with clear margins and they surgically removed with consideration of cosmetics. In this report (to our knowledge), we describe the first co-incidence of BCC and neurofolicullar hamartoma. The morphological features of smaller lesion and also expression of S100 protein in neural element helped us to achieve the diagnosis of neurofolicullar hamartoma. Based on variable reports of S-100 protein expression in NFH, as well as its similarity to peripheral nerve tumors, further studies are needed to determine the classification and differentiation of NFH.

Conflict of Interests: No conflict of interest has been expressed by the authors.

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