Pigmented Neurofibroma Of The Skin: A Case Report And Review Of The Literature
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Citation

Abstract
Pigmented neurofibroma (PNF) of the skin is a rare variant of neurofibroma that is frequently seen among the patients with neurofibromatosis type 1, and it accounts for less than 1% of all neurofibromas. We are reporting such a case with a review of the pertinent literature to highlight the differential diagnosis.

SOURCE OF SUPPORT
None

INTRODUCTION
Pigmented neurofibroma (PNF) of skin is a rare variant of neurofibroma. It has often been confused with other pigmented cutaneous tumors, especially pigmented dermatofibrosarcoma protuberans. The purpose of this paper is to report a case of PNF, and to review its differential diagnosis.

CASE REPORT
A 69-year-old female presented a 0.5 cm flat, pigmented skin lesion on the right dorsal forearm present for many years. She did not have any other skin lesions or any evidence of neurofibromatosis. The past medical history and the family history were unremarkable.

Microscopic examination of the biopsied skin lesion showed a normal epidermis with a pigmented spindle cell proliferation in the dermis (Figure 1A).

The tumor cells were spindled and elongated with wavy pink-blue cytoplasm within myxocollagenous matrix (Figure 1B).

The nuclei were oval, spindle or comma shaped with the finely dispersed chromatin. The nucleoli were inconspicuous. No mitotic figures are identified. Scattered cells with coarse granular dark-brown pigments in the cytoplasm were present in the tumor (Figure 1B inset). Immunostaining showed that the tumor cells were diffusely positive for Vimentin, S-100 protein, and Melanin-A.
Figure 2
Figure 1B: H&E, 20X. Tumor cells are loosely arranged in an abundant myxocollagenous matrix. The inset shows spindle tumor cells with coarse granular dark-brown pigments in the cytoplasm.

Figure 3
Figure 1C: Immunostain, Melanin A, 40X. Positive spindle cells.

Figure 4
Figure 1D: Immunostain, S-100 protein, 40X. Positive spindle cells.

DISCUSSION
Pigmented neurofibroma (PNF) \[1, 3, 4, 5\], also named melanotic neurofibroma, is a rare variant of neurofibroma. PNF is derived from neural tissue, and it has a typical histological appearance of a neurofibroma with scattered melanin-laden cells. Immunohistochemical studies demonstrate that the non-pigmented spindle cells are positive for S-100 protein, Melan-A, and CD34, but negative for HMA-45. But the melanin-producing cells are S-100 (+), MITF (+), Melan-A (+) and HMB45 (+/-), but CD34 (-) \[3\].

The differential diagnosis of PNF includes storiform neurofibroma, melanotic schwannoma, and cellular blue nevus. PNF is frequently confused with storiform neurofibroma, also called pigmented dermatofibrosarcoma protuberans (pigmented DFSP) or Bednar tumor \[2\]. DFSP exhibits an extensive storiform growth, has greater immunoreactivity for CD34, and lacks a diffuse S-100 staining. In DFSP, only pigmented cells have S-100 expression, and non-pigmented spindle cells lack expression of S-100, HMB-45 and Melan-A, although they are CD34 positive. DFSP is not associated with neurofibromatosis, and has a high local recurrence rate, while PNF is clinically non-aggressive. The dendritic pigment cells in DFSP are randomly distributed, while the pigmented cells in PNF tend to localize in the deep dermis and subcutis. Cellular blue nevus typically occurs in young adults. The tumor has prominent nested growth pattern and consists of pigmented dendritic cells and spindle cells. Melanotic schwannoma is a rare tumor linked to the autosomal dominant Carney’s disease. It is usually located deeply, and the pigmented cells
in this tumor are larger with an epithelioid appearance. The Schwann cells are S-100 (+), MITF (-), Melan-A (-), tyrosinase (-) and HMB45 (-)

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References
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