Violaceous Papule With an Erythematous Rim

Ania Henning, MD; Matthew Powell, MD; Tammie C. Ferringer, MD

A 35-year-old man presented with a reddish brown papule on the left upper chest of 1 year’s duration that had changed color to reddish purple. Physical examination revealed a 6-mm violaceous papule with an erythematous rim.

THE BEST DIAGNOSIS IS:
- a. angiokeratoma
- b. angiosarcoma
- c. Kaposi sarcoma
- d. targetoid hemosiderotic hemangioma
- e. verrucous hemangioma

H&E, original magnification ×100.
THE DIAGNOSIS:
Targetoid Hemosiderotic Hemangioma

Targetoid hemosiderotic hemangioma (THH), also known as hobnail hemangioma, is a benign vascular tumor that usually occurs in young or middle-aged adults. It most commonly presents on the extremities or trunk as an isolated red-brown plaque or papule. Histologically, THH is characterized by superficial dilated ectatic vessels with underlying proliferating vascular channels lined by plump hobnail endothelial cells. Targetoid hemosiderotic hemangioma typically involves the dermis and spares the subcutis. The vascular channels may contain erythrocytes as well as pale eosinophilic lymph, as seen in our patient (quiz image). The deeper dermis contains vascular spaces that are more angulated and smaller and appear to be dissecting through the collagen bundles or collapsed. A variable amount of hemosiderin deposition and extravasated erythrocytes are seen. Histologic features evolve with the age of the lesion. Increasing amounts of hemosiderin deposition and erythrocyte extravasation may correspond histologically to the recent clinical color change reported by the patient.

Verrucous hemangioma is a rare congenital vascular abnormality that is characterized by dilated vessels in the papillary dermis along with acanthosis, hyperkeratosis, and irregular papillomatosis, as seen in angiokeratoma. However, the vascular proliferation composed of variably sized, thin-walled capillaries extends into the deep dermis as well as the subcutis (Figure 1). Verrucous hemangioma most commonly is reported on the legs and generally starts as a violaceous patch that progresses into a hyperkeratotic verrucous plaque or nodule.

Angiokeratoma is characterized by superficial vascular ectasia of the papillary dermis in association with overlying acanthosis, hyperkeratosis, and rete elongation. The dilated vascular spaces appear encircled by the epidermis (Figure 2). Intravascular thrombosis can be seen within the ectatic vessels. In contrast to verrucous hemangioma, angiokeratoma is limited to the papillary dermis. Therefore, obtaining a biopsy of sufficient depth is necessary for differentiation. There are 5 clinical presentations of angiokeratoma: sporadic, angiokeratoma of Mibelli, angiokeratoma of Fordyce, angiokeratoma circumscription, and angiokeratoma corporis diffusum (Fabry disease). Angiokeratomas may present on the lower extremities, tongue, trunk, and scrotum as hyperkeratotic, dark red to purple or black papules.

There are 3 clinical stages of Kaposi sarcoma: patch, plaque, and nodular stages. The patch stage is characterized histologically by vascular channels that dissect through the dermis and extend around native vessels (the promontory sign) (Figure 3). These features can show

FIGURE 1. Verrucous hemangioma. A proliferation of dilated vascular spaces filling the papillary dermis and extending deep into the reticular dermis and subcutaneous adipose tissue (H&E, original magnification ×20).

FIGURE 2. Angiokeratoma. Dilated vascular spaces within the papillary dermis of an acanthotic epidermis with hyperkeratosis (H&E, original magnification ×100).

FIGURE 3. Kaposi sarcoma. Slitlike dilated vascular channels dissecting through reticular dermal collagen and around native vessels (promontory sign) (H&E, original magnification ×200).
Angiosarcoma is a malignant endothelial tumor of soft tissue, skin, bone, and visceral organs.11,12 Clinically, cutaneous angiosarcoma can present in a variety of ways, including single or multiple bluish red lesions that can ulcerate or bleed; violaceous nodules or plaques; and hematomalike lesions that can mimic epithelioid neoplasms including squamous cell carcinoma, basal cell carcinoma, and malignant melanoma.11,13,14 The cutaneous lesions most commonly occur on sun-exposed skin, particularly on the face and scalp.12 Other clinical variants that are important to recognize are postradiation angiosarcoma, characterized by MYC gene amplification, and lymphedema-associated angiosarcoma (Stewart-Treves syndrome). Angiosarcoma can have a variety of morphologic features, ranging from well to poorly differentiated. Classically, angiosarcoma is characterized by infiltrating vascular spaces lined by atypical endothelial cells (Figure 4). Poorly differentiated angiosarcoma can demonstrate spindle, epithelioid, or polygonal cells with increased mitotic activity, pleomorphism, and irregular vascular spaces.11 Endothelial markers such as ERG (erythroblast transformation specific–related gene) (nuclear) and CD31 (membranous) can be used to aid in the diagnosis of a poorly differentiated lesion. Epithelioid angiosarcoma also occasionally stains with cytokeratins.13,14

REFERENCES


FIGURE 4. Angiosarcoma. Vascular spaces lined by hyperchromatic and markedly atypical endothelial cells dissecting through the collagen (H&E, original magnification ×200).