A Rare Case of Arteriovenous Hemangioma Clinically Mimicking Pigmented Nevus

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Abstract

Arteriovenous hemangioma (AH) is a rare benign vascular skin lesion, which typically appears in the skin of the face and extremities and most commonly occurring on the head and neck region with appearances as single, raised, red, or violaceous papules; sometime tan papule. A case of AH clinically mimicking pigmented nevus in 19-year-old woman was reported. This is the first case in Dermatology Department of Dr. M. Djamil Padang Hospital. She complained about a black pimple on the right lower arm since one month. Physical examination: there is a solitary black papule, with 0,3x0,4 mm, round shape, fine scales, well defined, regular border with irregular surface. Histopathology findings: the lesions consist of thicked-walled and very dilated thin-walled vessels that full-filled with erythrocytes and are lined by an endothelial layer that suitable for AH. Arteriovenous hemangioma is a tumor of middle-age to elderly adults with a peak incidence in the fourth and fifth decades of life. In this case, the patient was young adult and clinically the lesion mimicking pigmented nevus.

Keywords: arteriovenous hemangioma, rare case, pigmented nevus

INTRODUCTION

Arteriovenous hemangioma (AH), also known as cirsoid aneurysm or acral arteriovenous tumor, represents a benign vascular skin lesion, which typically appears in the skin of the face and extremities.¹ ² It is a tumor of middle-age to elderly adults with a peak incidence in the fourth and fifth decades of life. Biberstein and Jessner reported the first case of AH in 1956 as “cirsoid aneurysm”. The entity then lay relatively dormant in the literature until 1974 when Girard et al. characterized the salient clinical and histologically similar lesions that they termed arteriovenous hemangioma. Subsequent
reports by Carapeto et al. and Connely and Winkelmann have confirmed the benign nature of these vascular tumors and have emphasized the acral pattern of distribution by the assignment of the name acral arteriovenous tumor.1,2

Arteriovenous hemangioma maybe cutaneous or mucosal. These lesions grow slowly and achieve only a small size with range from 1mm to 3 cm, with an average of 4 to 6 mm in months to many years. Most are asymptomatic, but a minority of patient complain enlargement, pain, or pruritus.3 The lesions most commonly present as single, raised, red, or violaceous papules; sometime tan papule on the head or neck. Microscopically, AH lesions are well-circumscribed but uncapsulated. They are composed of an intimate admixture of thick-walled and thin-walled blood vessels distributed within the superficial and middle dermis.1 Muscular walled vessels in haphazard cluster in the dermis. More vessels have features of veins than of arteries (and lack internal elastic lamina with an elastic stain).3

Enzinger and Weiss divide arteriovenous hemangioma into two types, "deep" and "superficial". The so-called “deep” type, is associated with varying degrees of shunting, and is regarded as a malformation. It is generally affect young adult and adolescent. Although called “deep”, lesions can occur close to the skin and they may pulsate or writhe due to afferent arterial blood flow if large shunts are present. These lesions are referred as cirrhotic or arteriovenous aneurysms. The superficial form includes lesions classified as arteriovenous haemangioma or acral arteriovenous tumor by Capareto et al. and generally affected middle age to elderly adult.2,4

The pathologic nature and etiology of Ahs remain a matter of speculation, and both congenital and acquired Ahs have been reported.5 However, hamartomatous proliferation either of the subpapillary vascular plexus or of the Sucquet-Hoyer canal of the true glomous is proposed as a possible histogenetic mechanism. Treatment of choice is complete surgical excision.6

CASE REPORT
A 19-year-old woman came to Dermato-Venereology out-patient Department of Dr. M. Djamil Hospital on December 28th 2011 with chief complaint there was a black pimple on the right lower arm since one month ago. Initially, about 2 years ago, there was a tiny reddish pimple arised on her right lower arm. The size of the pimple only as pinhead size. The pimple was painless, sometimes itchy, not easy to bleed although she scratched it, and not become large since appeared. In last two months, she often touched the pimple and scratched it because itchy. Since one month ago, she felt the pimple become larger and easily bleed after she scratched it and the color turned into black. On physical examination we found a solitare black papule, with 0,3x0,4 mm in diameter, round shape, fine scales, the regular border with irregular surface. Then, we diagnosed this patient with pigmented naevus at the beginning with differential diagnosis pigmented basal cell carcinoma (T1 N0 M0) and pyogenic granuloma. To exclude those differential diagnosis, we did histopathology examination by skin biopsy and the result showed mild acanthosis dan hyperkeratosis on epidermis and superficial dermis containing thicked-walled and very dilated thin-walled vessels that full-filled with erythrocytes and are lined by an endothelial layer. There are also seen bleeding area and mild lymphocytes perivasculer. This features supported to AH.

Figure 1. Black pimple on the right lower arm
DISCUSSION

Arteriovenous hemangioma (AH) is a rare vascular anomaly, with the principal histologic feature of which is a collection of thick-walled and thin-walled blood vessels with the structural characteristics of both arteries and veins. The exact incidence of AH is unknown. In most series, males are predominant.²,⁴

We reported a case of superficial arteriovenous hemangioma in a 19-year-old woman. The problems in this case are: a rare case and diagnosis. There are two forms of AHS according to Enzinger and Weiss, superficial and deep AH. Despite benign histology, deep lesions may produce serious systemic signs and symptoms such as heart failure and Kasabach-Merritt syndrome, because of extensive arteriovenous shunting and soft tissue hypertrophy. Diagnosis is usually made in conjunction with clinical, arteriographic, and radiographic investigations. By contrast, superficial AHs do not have systemic implications, but may nevertheless prove troublesome because of their tendency to affect the skin of the face, particularly the lips and perioral area. Symptoms are mild and include pain and intermittent bleeding.²,⁴

In this case, the patient was a young lady, and she did not have any systemic signs and symptoms. Mostly symptoms that she complained in last one month were enlargement and discoloration of the lesion, pruritus and intermittent bleeding.

Initially, we diagnosed this patient as pigmented nevus (common acquired nevomelanocytic nevus) because at the first time the patient came into hospital, the lesion was a small blackish papule that mimicking nevus. Pigmented nevus or acquired nevi primarily develop during childhood and early adulthood. Common acquired nevi vary considerably in their gross appearance. In general, appearance to the naked eye is orderly: the lesions have a homogeneous surface and coloration pattern, round or oval shape, regular outlines, and relatively sharp borders. Common acquired nevi may be papillomatous, dome-shaped, pedunculated, or flat-topped and usually are flesh-colored, pink, or brown. Very dark brown and black are unusual colors for common acquired nevi in lightly pigmented people. In contrast, dark pigmentation is usual for common acquired nevi in people who have darkly pigmented skin. The surfaces of nevi may reveal hair that is less than, equal to, or greater than that of surrounding skin. Based on histopathology features, nevomelanocytes in the epidermis have a nuclear size similar to or larger than nuclei of epidermal melanocytes. Nevomelanocytes are arranged in nests surrounded by a smooth perimeter of epidermis and separated from nevomelanocytes by a retraction artifact.⁷

The second possibility was pigmented basal cell carcinoma (BCC). From anamnesis there was reddish papule in right lower arm that turned into black in color (hyperpigmented) and become larger in last one month. The lesion also easily bleeding when the patient scratch it. Pigmented BCC is sub-type nodular BCC that exhibits increased melanization. Pigmented BCC shows histologic features similar to those of nodular BCC (characterized by nodules of large basophilic cells and stromal retraction) but with the addition of melanin. Approximately 75% of BCCs contain melanocytes, but only 25% contain large amounts of melanin. The melanocytes are interspersed between tumor cells and contain numerous melanin granules in their cytoplasm and dendrites. Although the tumor cells contain little melanin, numerous melanophages populate the stroma surrounding the tumor.⁸

The third possibility was pyogenic granuloma (PG), also known by its correct histopathologic description lobular capillary hemangioma. It is one of the most common vascular tumors of infants, children, and can also occur in adults, particularly in young
adults and also pregnant women. Pyogenic granuloma usually presents as a solitary, red, rapidly growing papule or nodule, often with a subtle collarette of scale. Typical locations include the cheek or forehead but virtually any body site may be affected. They often develop an eroded surface, with subsequent bleeding which can be profuse, resulting in the moniker the band-aid disease. The lesion of PG does not involute spontaneously, but simple curettage with electrocautery is usually curative.

Microscopically, this is an exophytic, often ulcerated lesion characterized by lobulated proliferation of capillary sized vessels in a loose and oedematous stroma. The surface epithelium is attenuated and at the margin an epidermal collarette is formed by elongated rete ridges. Clinically, the lesion of this patient can mimick PG as papule with fine scales and easily bleed.

The definite diagnosis in this patient was superficial AH. This diagnosis was made based on histopathology examination. There are some differential diagnosis in this case, but all of that can exclude by histopathology finding. The lesion was started as a single tiny reddish pimple arised on her right lower arm since two years ago. The lesion asymptomatic, grow slowly and achieve only a small size. We did biopsy excision and the result of histopathology examination was the lesions consist of thicked-walled and very dilated thin-walled vessels that full-filled with erythrocytes and are lined by an endothelial layer that suitable for AH.

Since the first case report in 1956, 4 series studies documenting 69, 15, 47, and 6 cases have demonstrated the clinicopathologic features of AH. Beside face and neck region, the skin of the limbs is the next most commonly affected location. Regardless of anatomic site, affected patients have a wide age range, with the fifth and sixth decades most commonly affected.

Barret Aw et al. (London, 2000) reported that from 470 vascular lesions between 1952 and 2000, 36 lesions (7.7%) from 35 patients had the features of superficial AH. All were situated intraorally or on the lip vermilion. Akiyama M et al. (Japan,2001) also reported 4 cases of AH in chronic liver disease. All patients showed an asymptomatic solitary dome-shaped reddish papule, 5±10 mm in diameter, on the face (first, second and third patients) or chest (fourth patient). Histopathological examination of the tumours revealed features of AH, namely a well-circumscribed lesion composed of vascular structures of various sizes reminiscent of arteries and veins.

The origin, cause, and nomenclature of AH have been debated at length, confusion often centering around the terms hemangioma and vascular malformation. Some authorities stipulate that hemangiomas occur during the first month of life and proliferate rapidly to a plateau, after which they stabilize and gradually involute. Endothelial or mast cell hyperplasia can be prominent. By contrast, vascular malformations are present at birth, grow in proportion to somatic growth, and never regress spontaneously. The blood vessels are abnormal arteries, veins, capillaries, or lymphatics and rarely show endothelial or mast cell hyperplasia. Based on these criteria, however, superficial AH may have the features of both entities.

There was no case of superficial AH with lesion clinically mimicking pigmented nevus has been reported. This is the first case of AH in our department. In this case, we did biopsy excision with concideration that the lesion was small, occurring of discoloration and bleeding of the lesion that tend to be a carcinoma and to exclude the differential diagnosis. The choice of treatment of AH is surgical excision, and recurrences have never been reported. The prognosis this lesion was bonam.

REFERENCES:


