CASE REPORT

Capillary haemangioma on the palate: a diagnostic conundrum

Pooja Singh,1 Anuj Singh Parihar,2 Sana Noor Siddique,3 Pooja Khare4

SUMMARY

Haemangiomas are benign tumours of blood vessel origin and are classified as capillary, cavernous or central. They appear as flat or raised reddish-blue lesions and are generally solitary, affecting women in younger age groups. The tumour may be slowly progressive, involving extensive portions of the superficial and deep blood vessels, and affect function, depending on location. They are common in the head and neck region but rarely in the oral cavity. Oral lesions generally appear on the lips, buccal mucosa and tongue, but rarely on the palate. As the lesion can be confused with pyogenic granuloma, histopathological examination is important for a final diagnosis. The case presented here signifies a rare location of a capillary haemangioma on the palate in a middle-aged man. The lesion was diagnosed by histopathology after surgical excision.

BACKGROUND

The term haemangioma is a generic one used to describe congenital hamartomas and vascular malformations. In general, haemangiomas are developmental and are often recognised at an early stage. They are considered to be the most common tumour of childhood, occurring in about 5–10% of children <1 year of age, exhibiting a rapid growth phase with endothelial cell proliferation followed by gradual involution. There is however an additional neoplastic form that appears in middle life or in older people. In the oral cavity they may occur in any area, at any age, without any racial or gender predilection. However, more than 50% are found in patients >40 years of age. The lips, buccal mucosa and tongue are the most common sites of occurrence, while the hard and soft palates and uvula are rarely affected. There occurrence on the palatal mucosa is extremely rare. The present case is of importance in view of the involvement of the hard palate, an extremely rare site (ie, hard palate involvement in an older age group).

CASE PRESENTATION

A 31-year-old man presented with a growth on the left anterolateral part of the hard palate of 3 years’ duration with a complaint of discomfort during mastication and speech. The growth was the size of a peanut when he first noticed it 3 years previously and had grown to the present size. The growth was asymptomatic; there was no history of pain, pus discharge or bleeding, or any other secondary symptoms. His medical dental and family history was non-contributory.

INVESTIGATIONS

Intraoral soft tissue examination revealed a solitary broad-based spherical pink growth with a well-defined border and pebbly surface. It was present on the left anterolateral part of the hard palate just 2 cm lateral to the midline on the left side in an area between the maxillary first and second premolars, measuring about 1.5 cm in diameter. On palpation it was non-tender, soft to firm in consistency, compressible but not reducible and non-pulsatile. The patient’s oral hygiene status was fair and no other relevant intraoral findings were evident (figure 1).

In the clinical context of a well-demarcated slow-growing lesion, the differential diagnosis included congenital hamartomas, pyogenic granuloma and other vascular malformations. The lesion was surgically excised. The histopathological examination confirmed it to be a capillary haemangioma, with endothelial cell proliferation and formation of capillary blood vessels.

Figure 1 Intraoral view.

Unusual presentation of more common disease/injury

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Unusual presentation of more common disease/injury

The complete haemogram was within normal limits (figure 2).

The histopathological examination of the excised mass showed parakeratinised stratified squamous epithelium of varying thickness (figure 3). The fibrous connective tissue showed numerous endothelial lined blood vessels of varied sizes and few blood vessels, which were yet to be lumened. Numerous mixed inflammatory components were present (figure 4). Histopathological features were suggestive of capillary haemangioma with an inflammatory component. The inflammatory component in histopathological examination can be due to trauma. Hence on correlation of the clinical and histological findings, a final diagnosis of capillary haemangioma was established.

DIFFERENTIAL DIAGNOSIS

The differential diagnosis of peripheral ossifying fibroma was made due to the similar clinical presentation of the lesion. However, the peripheral ossifying fibroma shows resorption of bone at the site of the lesion and radiographs are advised to rule out bony destruction, but no such finding was detected in the intraoral radiograph taken. Another similar appearing lesion taken into consideration was irritational fibroma, as the lesion appeared to be red to purple with a pebbly surface and on palpation it was compressible.

TREATMENT

As the lesion was small in size, an excisional biopsy was done under local anaesthesia. There was excessive bleeding at the time of biopsy, which was controlled after application of pressure packs and local styptics.

OUTCOME AND FOLLOW-UP

Wound healing was uneventful and no recurrence was noted after 5 months of follow-up.

DISCUSSION

Haemangiomas, whether they represent a true neoplasm, malformation or hamartomas, have been the subject of continuous debate, and use of the terms haemangioma and malformation can be confusing. Most pathologists separate haemangioma, composed primarily of blood vessels of similar size and type, from malformations, composed primarily of arteries and veins of varying sizes. In 1982, Mulliken and Glowaki developed a classification of vascular tumours based on endothelial characteristics.

Haemangiomas of the oral cavity may exist as small or large superficial growths with varying degrees of penetration into soft tissues or as monstrous growths extending to the oesophagus. They may also be multicentric with a cobblestone appearance. Most of the small superficial lesions, especially pedunculated ones, tend to be capillary haemangiomas, whereas the large superficial or deep lesions tend to be of the cavernous or mixed variety.

Capillary haemangiomas can be sessile or pedunculated, soft, smooth or irregular, and bulbous in outline, and are painless unless traumatised. They vary in colour from deep red to purple and blanch on application of pressure. Those on the palatal mucosa are solely of the capillary type. In the present case, the lesion was broad based, red to purple with a pebbly surface. Periodontally they may appear to arise from the mediolateral gingival papillae and spread laterally to involve adjacent teeth. However, in the present case, the lesion was not attached to the gingiva and was 1 cm medial to it.

This varied appearance, infrequent association with palatal gingival and tendency to mimic other lesions leads to confusion with other clinical entities, such as pyogenic granuloma, chronic inflammatory gingival hyperplasia, epulis granulomatosa, telegenesis, angiosarcoma or squamous cell carcinoma. In the present case, with the clinical appearance, a working diagnosis of pyogenic granuloma was assumed; however, histopathological examination confirmed a diagnosis of capillary haemangioma.

The usual capillary haemangioma is composed of many small capillaries lined by a single layer of endothelial cells, supported by a connective tissue stroma of varying density. It bears considerable resemblance to the young granulation tissue and is mostly identical to some cases of pyogenic granuloma. Some cases show remarkable endothelial cell proliferation. Management is based on several factors, such as age of the patient, size, extent and clinical characteristics of the lesion. The smaller lesions can be carefully and successfully excised, as in the present case. Most small capillary haemangiomas reported...
in the literature have been treated with excision or curettage.\textsuperscript{10} Local complications, such as bleeding or postoperative ulcerations, tend to occur. During excision in the present case, bleeding was controlled with application of pressure and/or styptics, while wound healing was uneventful in our case.

Twitter Follow Anuj Parihar at @anujhimself

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REFERENCES


