Angina Bullosa Hemorrhagica- A Clinical Report

Dipti Singh¹, Sudhanshu Agrawal², M C Shashikanth³, Deepak U³ and Anshul Mehra³

ABSTRACT

Aim: To report a case of 50 year old female patient with angina bullosa hemorrhagica on the right buccal mucosa since 15 days.

Summary: Angina Bullosa Hemorrhagica is the term used to describe benign subepithelial oral mucosal blisters filled with blood that are not attributable to a systemic disorder or haemostatic defect. It is a very rare condition. Elderly patients are usually affected and lesions heal spontaneously without scarring. The pathogenesis is unknown, although it may be a multifactorial phenomenon. Trauma seems to be the major provoking factor and long term use of steroid inhalers has also been implicated in the disease. We present a 50-year-old patient with angina bullosa hemorrhagica. Trauma by sharp cusp of adjacent tooth and metal crown were identified as etiological factors in this case. Lesions healed after removal of metal crown and rounding of cusp.

INTRODUCTION

In 1967, Badham coined a new term, angina bullosa hemorrhagica (ABH), to describe oral blood-filled vesicles or bullae that could not be attributed to a blood dyscrasia, vesiculo-bullous disorders, systemic disease or other known causes.¹ It is a disorder characterized by the acute formation of a blood-filled blister in the oral and oropharyngeal mucosa. Lesions of ABH occur mainly on the soft palate. Elderly patients are usually affected.

ABH is more common than previously suggested and, during the past decade, several studies have been published. In 1933, Balina of Argentina had already described the same lesions under the term traumatic oral hemophlyctenosis (TOH). He also postulated a trauma-induced origin, especially in patients with senile capillary changes. In 1969, 14 patients were presented, and, in 1976, the clinical and histologic features were detailed and documented.² This entity was then named recurrent oral hemophlyctenosis (ROH). As Kirtschig and Happel pointed out, the term ABH is misleading because most bullae arise in the oral cavity and are not consistent with lesions usually called “angina”; they proposed a more appropriate name for the disease: stomatopompholyx hemorrhagica. The authors believe that Balina was the first to describe this condition and suggested the use of the name ROH.³ The aim of this article is to report a new case of ABH, in an attempt to distinguish it from other blood containing bullae of the oral mucosa and to describe its management.

CASE REPORT

A 50-year-old female patient came to the Department of Oral Medicine & Radiology with a chief complaint of recurrent oral blisters on the right buccal mucosa recorded in a span of 15 days (Fig. 1). Pain and burning sensation was also present in that region. Intraoral hard tissue examination there was generalized attrition. Metal crown was present in relation to maxillaryright second molar (17) and third molar (18), which were tender on percussion and were grade I mobile. Sharp cusps were present in relation to maxillary right first (16), second (17) and third molar (18) and mandibular right first (46), second (47) and third molar (48).

Figure 1: Extraoral photograph of patient
In soft tissue examination the gingiva was inflamed and three blood filled bullae were present on the right buccal mucosa approximately 3 x 3 mm which were smooth surfaced, red in color and surrounding area was erythematous (Fig. 2). On palpation they were tender and bleed on manipulation. The patient reported no blood dyscrasias, anticoagulant therapy, or liver disease and was generally having good health. Other than these oral blood blisters, the patient reported no oral conditions and no skin or eye lesions. Family history was also negative. Biopsy was advised but patient was not willing.

So, based on history of continuous trauma from teeth to the mucosa and clinical examination we came to a provisional diagnosis of ABH. Differential diagnosis was made to exclude other mucosal or cutaneous diseases such as erythema multiforme, bullous lichen planus, pemphigus, pemphigoid and epidermolysis bullosa. Hematological blood cell count and differential and prothrombin time (PT) and activated partial thromboplastin time (APTT) tests were carried out and the findings were within normal limits.

In treatment removal of metal crown and grinding of sharp cusps was done. Ointment Mucopain (Benzocain20%), Tantum oral rinse (Benzydamine hydrochloride) was prescribed and patient was recalled after 10 days. Patient came after 10 days and the lesion was healed (Fig 3).

DISCUSSION

ABH, first described by Badham in 1967, is a condition characterized by the rapid formation of a blood-filled blister on the oral mucosa. The disorder is now considered to be more common than the literature or conventional wisdom previously suggested. ABH mainly affects the soft palate, but lesions also can develop on other oral sites including the buccal mucosa, lip, and the lateral surface of the tongue; masticatory mucosa of the hard palate and gingiva does not seem to be affected. ABH patients have been mainly middle-aged and elderly; lesions have not been documented in children less than 10 years of age. There is no apparent gender predilection. ABH has been considered as an idiopathic condition. The onset is sudden and minor mucosal insults may be involved in the pathogenesis. It may also follow trauma caused by eating, hot drinks, dental procedures or shouting. It is also noteworthy that mastication significantly increases the blood flow rate in the soft palate via parasympathetic reflex vasodilatation and hard or crispy may injure the palate which leads to ABH. The use of steroid inhalers in asthmatic patients is a possible aetiologcal factor. In the largest published series of 30 patients, no precipitating factor was found in 47%. Hosain and colleagues reported a case of postoperative ABH caused by intubation and extubation, describing a patient with a single blister at the junction of the soft and hard palate that did not compromise the patient’s airway. Lesions predominantly occur on the soft palate. The intact bulla is red to purple in color. Blisters usually reach 2-3 cm in diameter and burst spontaneously, leaving ragged ulcers that heal without scarring but the lesions can recur.

The diagnosis of ABH is largely clinical, and includes the elimination of other disease process at histology. Histopathological features of ABH include parakeratotic epithelium with a subepithelial separation from the underlying lamina propria. Superficially located vesicles filled with erythrocytes and fibrins are seen. The inflammatory cell infiltrate, when present, consists primarily of lymphocytes. Neutrophils and eosinophils seen in other blistering disorders are not present. Immunoflorescence demonstrates no evidence of IgG, IgM, IgA or C3 antibodies within the epithelium or basement membrane zone.

Lesions of ABH can be easily confused with those occurring in many dermatological and systemic disorders. Even if typical history of rapid blistering disease if there is absence of any dermatological, hematological or systemic sign and normal healing of the ulcers generally lead to the ABH diagnosis.
Patients with bleeding disorders (thrombocytopenia and von Willebrand’s disease) can present with intra-oral blood filled lesions but hemostatic function test will distinguish these conditions. The absence of desquamative gingivitis and nasal or conjunctival mucosal involvement will differentiate it from benign mucous membrane pemphigoid. Linear IgA disease and dermatitis herpetiformis usually can be differentiated by the presence of a pruritic rash. In oral Bullous Lichen Planus bullae are often associated with striated pattern. The target like lesion of the skin in erythema multiforme help to distinguish it. The hemorrhagic bullae found in amyloidosis are usually persistent and other clinical features include macroglossia and petechiae. Epidermolysis Bullosa can be differentiated by the presence of bullous skin lesions.

The management of a patient presenting with oral blood-filled bullae should start with detailed medical history and careful examination to differentiate ABH from other more serious diseases. The lesion should be biopsied to perform histology and direct immunofluorescence in order to exclude more serious diseases. A complete blood count and a baseline coagulation tests should always be performed to exclude blood disorders. The patient should be reassured of the benign nature of the blisters. Large palatal or pharyngeal blister causing a choking sensation should be surgically treated if still intact. Management of these lesions should be symptomatic. Long term follow up is recommended to positively exclude other conditions which may present with oral blood-containing bullae.

CONCLUSION

The diagnosis is difficult in patients as ABH is asymptomatic and heals spontaneously without scarring and its rare appearance. The diagnosis of the lesion is very important as a rapidly expanding blood-filled bulla in the oropharynx can cause upper airway obstruction. Therefore, a high level of suspicion is warranted on part of dentists who may the first to encounter the lesion.

REFERENCES