Oral Manifestations of a Patient with Epidermolysis Bullosa: Case Report

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Submission: June 27, 2017; Published: December 13, 2017

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Abstract
Epidermolysis bullosa acquisita (EBA) is a chronic autoimmune bullous disease characterized by the presence of IgG and IgM antibodies at the level of basement membrane. It is rare in humans and animals with an incidence ranging from 0.2 to 0.5 new cases per million and per year. This dermatological condition is a severe autoimmune disease. Scarring of the extensor surfaces of the extremities, hands and feet are typical; milia occur frequently; and nails often become thick and dystrophic or are lost. The disorder affects both sexes equally and occurs in all racial and ethnic groups.

Keywords: Adherent fingers; Absent nails; Microstomia; Bullae; Preventive care

Introduction
Epidermolysis bullosa (EB) is a group of rare inherited disorders, usually detected at birth or early childhood [1-4]. Köhner coined the term ‘epidermolysis bullosa’ in 1886, but even before this time, Legg and Brocq had already provided a clinical description of the disease. Epidermolysis bullosa characterized by extreme fragility of the skin and mucous membranes, which gives rise to the formation of blisters following minor trauma [5]. This dermatological condition is a severe autoimmune disease [6,7]. Scarring of the extensor surfaces of the extremities, hands and feet are typical; milia occur frequently; and nails often become thick and dystrophic or are lost. The disorder affects both sexes equally and occurs in all racial and ethnic groups [8]. Epidermolysis bullosa has been classified into three major types depending upon the histological level of tissue separation [9]:

1. epidermolysis bullosa simplex is characterized by discontinuities in the epithelial keratinocyte layer;
2. junctionalepidermolysis bullosa involves separation within the basement membrane; and
3. dystrophicepidermolysis bullosa is characterized by discontinuities in the underlying connective tissue.

Each type of EB has various subtypes and these may vary in severity [10]. Skin biopsies are needed for appropriate diagnosis and classification for affected subjects.

Case Report

A 12-year-old female patient diagnosed with severe generalized junctional EB was referred to the Department of oral medicine and Radiology. The patient complaints of dental pain, halitosis, severe crustation of lips and limited mouth opening with ulcerations of buccal mucosa. The patient had one sister, aged 6 years old, who was unaffected by the disease. Her parents were also unaffected and were not consanguineous. Both sets of grandparents came from the nearby areas of same state. This type of illness had not previously appeared in the family. Physical examination revealed generalized worn-out skin, blistering and...
scar formation, with blisters and vesicles present especially on the head and neck. The patient’s few fingers were adherent, and her nails were absent (Figure 1,2). Scar formation had resulted in the formation of microstomia (Figure 3). The patient’s maximum mouth opening was 14 mm. Clinical examination showed multiple missing teeth, decay and poor oral hygiene (Figure 4), due in part to a soft diet and hand contractures.

The patient was a case of junctional epidermolysis bullosa which was confirmed by histopathology. Skin biopsy from fresh bulla showed a large dermo-epidermal bulla with mild chronic inflammatory infiltrate in dermis. Direct immune fluorescence showed IgG IgM deposition at the level of basement membrane, all the features were suggestive of junctional epidermolysis bullosa. The goal of treatment in the present case was to prevent blisters from forming and its resulting complications. Topical steroids and antibiotics were used to promote healing and prevent secondary infection of blisters. Since restoration of teeth was not possible due to severe crown destruction, extraction of few teeth was done in two sittings under local anaesthesia. The use of an aloe vera containing tooth gel at home was suggested to soothe the burning feeling affecting the gums. A mouthwash was also prescribed to help the salivary immune system protect the mucosal surfaces. In this case, minimal intervention has sofar preserved the oral cavity and monthly topic fluoride application helped to control dental caries. The patient maintains continuous contact with the health team to avoid complex treatments. Follow-up is advised after every 3 months for the evaluation of oral status.

Discussion

EB is a group of rare genetically determined disorders characterized by the development of blisters following minor or insignificant trauma or traction to the skin or mucosal surfaces. [11–13] Oral manifestations and dental involvement of EB vary in frequency, in severity and according to subtype. Dental management of individuals with EB has been reported previously by several authors [14-15]. Examinations have shown that the caries prevalence among individuals with dystrophic EB and functional EB is significantly higher than among healthy people [16]. Similar observations have been made with respect to the prevalence of plaque and gingivitis. However, the patients examined in these studies show no significantly higher prevalence of Candida albicans, Lactobacillus casei or Streptococcus mutans [17]. Wright et al. reported that none of their patients with EB showed any evidence of a decreased salivary flow rate [18]. Despite the severe cutaneous and extracutaneous involvement associated with inherited EB, the above authors found no evidence to support the hypothesis of abnormal salivary function or mucosal immunity in this disease. Taken together, these findings suggest that the rampant dental caries seen in the various forms of EB are more likely to be attributable to non salivary factors, such as enamel involvement, soft-tissue alterations and/or diet. Dental caries have a complex pathogenesis and may be influenced by a variety of genetic and environmental factors. Childhood EBA is a very rare disease and none of the previously described patients had dental abnormalities as a clinical manifestation of this disease. Interestingly, defects in enamel maturation and/or mineralization have been reported in dystrophic and junctional epidermolysis bullosa. The Col17-/- mice, a model to study non-Herlitz junctional epidermolysis bullosa enamel hypoplasia, exhibit imperfect amelogenesis demonstrated by malformed enamel rods and irregular enamel matrix [19]. Similar changes are observed in some cases of epidermolysis bullosa caused by mutations in laminin 5, a6b4 integrin or collagen VII [20]. The tooth abnormalities due to type VII collagen deficiency investigated in Col7a1-/- and COL7-rescued humanized mice may be attributed to poorly differentiated ameloblasts [21]. Defective...
enamel structure may provide a flourishing local environment for cariogenic bacteria by providing more adhesive and colonization potential and being more acid soluble. Hence, defective enamel appears to be a favoring factor for dental caries of childhood. Epidermolysis bullosa treatment is generally focused on support. Perforating the blisters contributes to accelerating the healing process and prevents continued lateral spread of the blisters. Currently, researchers are focusing their attention on gene and cell therapy, recombinant protein infusions, intradermal injections of allogenic fibroblasts and stem cell transplantation. Other developing therapies are directed toward the enhancement of wound healing and better quality of life for EB patients [22]. Some difficulties complicate the rehabilitation treatment of these patients, as the fragility of the mucosa and the microstomia. The lesions in the oral mucous membrane can be so severe that they interfere with nutrition as a result of the difficulty of food due to pain as in present case. Thus, the treatment of the EB patient must be modified in order to decrease the effects of local trauma to the tissues. Oral ulceration due to trauma during dental manipulation is unavoidable, but it could be limited by the mucosa lubrication before any manipulation with hydrocortisone cream, triamcinolone or petroleum jelly [23]. In addition, the use of small-sized instruments, short-shaft dental burs and hand pieces with a small-sized head is indicated. Dentures and restorations should be carefully adapted and highly polished; secondary infections should be prevented with the use of oral antiseptics or using topical antibiotics on existing bullous lesions. During the dental treatment in outpatient settings, the administration of local anaesthesia to patients with EB may also cause blister formation and should therefore be avoided whenever possible. Nevertheless, when necessary, local anaesthetic should be injected slowly and deeply into the tissues to prevent mechanical tissue separation and blistering. Owing to the difficulties related to the restorative dental treatment of these patients, the emphasis on the oral health management must be on prevention beginning at an early age. Regular application of topical and systemic fluoride, oral hygiene instructions and dietary guidelines minimize caries development and improves overall oral health.

Conclusion

Early dental management and preventive care to minimize caries development and to improve oral health is very important for patients with EB. Furthermore, they are often advised to reduce their consumption of cariogenic foods. Topical fluoride applications at regular intervals are also indicated. Ongoing patient monitoring must be stressed to minimize the need for restorative treatment in patients with EB.

References


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DOI: 10.19080/ADOH.2017.07.555704

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