Oral Medicine Case Book: Epidermolysis bullosa acquisita.

CASE REPORT
A 49-year-old female was referred to the Dermatology department to the Oral Medicine Department at the University of the Western Cape (UWC), Oral Health Centre, Tygerberg campus. The patient complained of a sore mouth and difficulty in brushing her teeth. She had been diagnosed with epidermolysis bullosa acquisita (classical type) 12 years ago and was being managed by her Dermatologist with topical and systemic steroids (Dovate® ointment and 10mg prednisone daily).

The extra oral examination revealed extensive sloughing of the skin of the hands, chest and back. The hands showed atrophic scarring, skin fragility and nail loss on numerous fingers, which also demonstrated restricted movement (Figures 1, 2).

The patient had limited mouth opening because of scarring related to repeated episodes of ulceration (Figure 3) and poor oral hygiene. Her gingiva was inflamed (Figure 4). The middle to anterior dorsal surface of her tongue was atrophic, smooth and erythematous, while the posterior dorsal tongue had white yellowish plaques (Figure 5). The orthopantomogram demonstrated multiple carious teeth with generalised, severe horizontal bone loss (Figure 6).

An appointment was scheduled for a scaling as well as extraction of root remnants and teeth with a hopeless prognosis. The patient was booked for a follow-up at the Oral Medicine and Periodontology Department two weeks later, but failed to return for her appointment.

DISCUSSION
Epidermolysis bullosa (EB) is a group of inherited blistering diseases, which can be present from birth, or be acquired and manifest in adult life.1 The first description of a patient with a bullous disease reminiscent of EB (with no associated familial history), was reported by Elliot in 1895.2 In the early 1970s, Roenigk et al. proposed the first diagnostic criteria for EBA.2

ACRONYMS
BMZ : basement membrane zone
DEJ : dermal-epithelial junction
DF : Direct immunofluorescence
EBA : Epidermolysis bullosa acquisita
EB : Epidermolysis bullosa
ELISA : enzyme linked immunosorbent assay.
IBD : inflammatory bowel disease
IF : Indirect immunofluorescence
SLE : systemic lupus erythematosus

Figure 1: Atrophic scarring and skin fragility on the back of the hands, nail loss and deformity of numerous fingers.

Figure 2: Severe sloughing and scarring, with restriction of movement.

The diagnostic criteria included:
1. Spontaneous or trauma-induced blisters resembling hereditary dystrophic EB
2. Adult onset of the disease
3. No associated family history for EB
4. Exclusion of all other bullous diseases2

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EBA is a rare, autoimmune, sporadic, subepithelial, mucocutaneous blistering disease, which usually occurs in adulthood. Skin fragility, non-inflammatory tense bullae, milia and extensive scarring typically characterize EBA. Otherwise, EBA may manifest as an inflammatory bullous eruption reminiscent of bullous pemphigoid or another sub-epithelial autoimmune blistering disorder.

Limited numbers of paediatric cases of EBA have been reported, but it is adults who are most commonly affected. Increased risks for EBA development based upon gender, ethnicity or geographic location have not been definitively established in the literature.

**Pathogenesis**

Although the pathogenesis of EBA is not clearly defined, evidence reveals that it involves an autoimmune process (type III hypersensitivity) with the production and deposition of antibodies against type VII collagen, which is a major constituent of the anchoring fibril (adhesion structures in the dermal-epidermal junction – DEJ). This theory concept is based upon several observations:

- Direct immunofluorescence (IF) microscopy of peri-lesional skin from patients with EBA showed antibody deposition at the basement membrane zone (BMZ).
- Indirect immunofluorescence (IF) revealed antibodies against type VII collagen in the serum of patients with EBA.
- Immuno-electron microscopy detected antibody deposition in the lamina densa area of the BMZ, a site consistent with the location of anchoring fibrils.
- Declining numbers of normal anchoring fibrils were present in the DEJ in EBA, a finding consistent with epidermal blistering and skin fragility.
- Animal models indicate that passive transfer of antibodies against type VII collagen can induce clinical features consistent with EBA.
- Serum levels of type VII collagen antibodies identified by ELISA correlate with the severity of skin lesions.
- Patients with bullous systemic lupus erythematosus (SLE), which is also accompanying with antibodies against type VII collagen, develop sub-epithelial blistering.

**CLINICAL VARIANTS**

Classical EBA manifestations were firstly defined as non-inflammatory, mucocutaneous bullae with accompanying skin fragility. EBA may also present as an inflammatory blistering disease characterized by descriptions that resemble bullous pemphigoid, linear IgA bullous dermatosis, mucous membrane (atrophic) pemphigoid, or Brunsting-Perry pemphigoid. The classical and bullous pemphigoid-like subtypes are the most common presentations of EBA.
Classical (non-inflammatory) EBA

Patients with classical EBA present with skin fragility and non-inflammatory tense vesicles and bullae, which rupture quickly and develop erosions. The parts that are frequently subject to minor trauma are the most common locations for lesion progression, such as the hands, feet, knees, elbows, and lower back. These blisters usually heal with scarring and milia (small epidermal inclusion cysts).1,5

Mucosal involvement is common in classical and also in inflammatory EBA. It can be subclinical with associated pain or itching, or may present as adhesions or erosions on the oral, nasal, pharyngeal, laryngeal, esophageal, ocular or anogenital mucosa.6

Nail loss, alopecia, fibrosis of fingers and hands and esophageal stenosis may manifest in severe cases.1,4

Inflammatory EBA

These subtypes of EBA are similar to other autoimmune sub-epithelial blistering disorders. Compared to classical EBA, skin fragility is not a characteristic feature.1

- **Bullous pemphigoid-like EBA**
  
  This disorder shares the clinical manifestation with bullous pemphigoid that is the most common autoimmune subepithelial blistering disease.1,4 Unlike the non-inflammatory lesions of classical EBA, the condition manifests with widespread tense bullae with inflamed or urticarial skin. The lesions are common in the trunk, extremities and skin folds similar to bullous pemphigoid. Scarring and milia are not obvious findings.5

- **Mucous membrane pemphigoid-like EBA**
  
  This may appear as a mucous-predominant disorder with clinical manifestations that are similar to mucous membrane pemphigoid.1,5 The initial disease findings are erosions and scarring on the mucosal surfaces of the mouth, upper esophagus, conjunctiva, anus or vagina.5

- **IgA bullous dermatosis-like EBA**
  
  This has clinical, histologic, and direct immunofluorescence (DIF) manifestations that resemble IgA-mediated bullous dermatoses.3 The clinical manifestations may present with the annular distribution of vesicles and bullae characteristic of linear IgA bullous dermatoses. Associated mucosal involvement is common.5,15

- **Brunsting-Perry pemphigoid-like EBA**
  
  The disorder resembles Brunsting-Perry pemphigoid, which is a rare sub-epithelial blistering disorder. It presents as vesiculobullous eruptions, primarily appearing on the head and neck and heals with scarring.1,5

**DIAGNOSIS**

EBA resembles other sub-epithelial blistering disorders in some clinical, pathologic and immunohistologic features. Establishing a diagnosis can be challenging. Once the initial assessment suggests the presence of an autoimmune sub-epithelial blistering disorder and reveals findings consistent with EBA, additional investigations may be utilized to verify the diagnosis.1

**Initial patient evaluation**

The evaluation should include the following:1

- A full patient history and complete skin examination which involves an evaluation of the morphology and distribution of skin lesions

This assessment assists in limiting the differential diagnosis. For example, a diagnosis of classical EBA must be considered in adults who present with consistent clinical manifestations. These include skin fragility and trauma-induced tense bullae, which result in milia and scars, with no family history of any hereditary blistering disorder. Because of the numerous EBA’s morphologies, the probability of EBA should still be considered when clinical descriptions indicate any other sub-epithelial blistering disease.

- A biopsy for routine histopathology and direct immunofluorescence (DIF)

A tissue biopsy of affected skin or mucosa must be taken for routine histologic examination to define the level of blistering. A peri-lesional skin or mucosal biopsy for DIF should also be obtained to identify autoantibody deposition. A punch biopsy of 4mm is classically used to retrieve tissue specimens.

- **Immunofluorescence at basement membrane zone (BMZ)-split skin**

Once the clinical, histologic and DIF results are reliable with an autoimmune sub-epithelial blistering disease, immunofluorescence on skin artificially split within the BMZ is helpful to rule out bullous pemphigoid and linear IgA bullous dermatosis, thus narrowing the differential diagnosis to EBA and a few other uncommon disorders.

Indirect immunofluorescence (IDF) microscopy on salt-split skin is a simple and reliable tool compared with DIF, which helps to subclassify subepidermal autoimmune bullous diseases into "roof" and "floor" binding conditions.1 EBA is a standard floor-binding sub-epidermal autoimmune bullous disease. On the other hand, the floor binding of salt-split skin is not only revealed in EBA but also in anti-laminin 332 mucous membrane pemphigoid and anti-p200 pemphigoid.7

EBA shares clinical manifestation with other multiple disorders. A list of these disorders, frequently mistaken with EBA, are listed below:15

- Bullous pemphigoid
- Linear IgA bullous dermatosis
- Porphyria cutanea tarda
- Bullous systemic lupus
- Recurrent dystrophic epidermolysis bullosa

**Management**

Treatment strategies are determined by the clinical presentation. However, a multidisciplinary approach is necessary, including a nutritionist, dermatologist, hematologist, plastic surgeon, ophthalmologist, cardiologist, gastroenterologist, dentist, nurse and an occupational therapist.1

The first line of treatment in most of EBA patients is colchicine because of its efficiency and less associated side effects.15 Colchicine is a familiar microtubule inhibitor, which also plays a role in regulating autoimmunity by inhibiting antigen presentation to T cells. Colchicines in high doses have been reported to be effective for both classical and inflammatory EBA patients.16 Diarrhoea is an adverse effect which limits its use in EBA patients with inflammatory bowel disease (IBD).1

Rituximab is a monoclonal antibody, which targets CD20 on both mature and immature B cells.15 As a result of destroying B cells, the circulating antibodies and B cells are reduced, leading to an increase in immunosuppression.8

Cyclosporine, an immunosuppressant, is often considered in the treatment of EBA.13 Studies have revealed that some EBA patients responded to cyclosporine use. However, patients require high doses (>1mg/kg) of the drug. Cyclosporine has long-term toxicity, but is a valuable treatment modality in patients who are non-responsive to EBA therapy.5

Systemic glucocorticoids in high doses have proven to be successful in the management of some EBA patients.15 Other immunosuppressants such as methotrexate, azathioprine, and cyclophosphamide may also be used.15 Prednisone and dapsone may also assist some EBA patients.6

Photopheresis was efficient in a cohort of EBA patients, including one in a critical condition. Photopheresis is used to treat mycosis fungoides and Sézary syndrome (which is a malignancy of skin-homing CD4+ lymphocytes).5 The drug is clinically described as erythroderma, lymphadenopathy, and blood involvement(15) and numerous autoimmune bullous disorders.5
Blister formation following mild mechanical trauma characterizes most of major types of EB and most EB patients may reveal systemic complications, such as genital, ocular and oropharyngeal infections with difficulty in swallowing.13

Implications for the oral health worker
EB patients need particular precautions during dental treatment due to the probability of soft tissue injury during their examination of the oral mucosa and skin.9 These patients are predisposed to dental caries as a result of their cariogenic diet; poor oral hygiene worsened by pain and limited mouth opening.11 Regular visits/ contact with the oral health team can help to avoid complex procedures and treatments.12

Several alternative treatments are often used as first aid therapies for blisterers. Aloe vera gel application decreases the sub-dermal temperature, affords a refreshed sensation, promotes antimicrobial activity and diminishes the healing period.13 Biotene® mouthwash minimizes blister formation by providing oral moisturizing and salvia stimulation, providing buffering capacity and antimicrobial activity.14

Nowadays, researchers are investigating treatments such as gene and cell therapy, intradermal injections of allogeneic fibroblasts, recombinant protein infusions and stem cell transplantation.15 Other developing treatments for EB patients are focusing on the improvement of wound healing and good quality of life.16

It is essential that all EB patients receive supportive care to decrease the possibility of skin trauma and to improve quality of life. This supportive care involves appropriate wound management and approaches for preventing trauma. Patients should be educated that harsh soaps, vigorous rubbing of skin during washing specifically with hot water, may aggravate the lesions and result in trauma. The application of sunscreen may reduce the exacerbation or stimulation of new lesions due to prolonged sun exposure. Finally, the patient should be well informed to recognize superinfections of the skin and to seek urgent medical care should they develop.

References