Bart Syndrome: a rare entity

Síndrome de Bart: uma entidade rara

Chitreddy Uma Reddy[a], Komar Suresh Reddy[b], Jaddu Jyothirmai Reddy[c]

[a] BDS, MDS, professor, Department of Oral Medicine & Radiology, Adesh Institute of Dental Sciences and Research, Barnala Road, Bathinda, Punjab - India.
[b] BDS, MDS, professor, Department of Oral and Maxillofacial Surgery, Manasarovar Dental College, Bhopal, Madhya Pradesh - India.
[c] BDS, MDS, senior lecturer, Department of Preventive and Community Dentistry, Darshan Dental College & Hospital, Ranakpur Road, Loyara, Udaipur, Rajasthan - India, e-mail: drjoe0802@gmail.com

Abstract

Introduction: Bart Syndrome is a rare inherited skin blistering disorder. It is also known as congenital transient mechano-bullous dermatosis and is one of the lesser known presentations of epidermolysis bullosa (EB). Case report: The objective of this report is to present a case of Bart Syndrome in a 3 day old newborn female baby. The skin lesions showed denuded areas with bullae rupturing easily to reveal painful eroded areas. Eroded lesions were distributed over the hands, feet, chest and on the face over the cheeks bilaterally. The lips were erythematous, eroded with tissue tags. Eroded, crustated lesions were seen on the labial mucosa and anterior palate. Histopathological examination revealed split localized to the epidermis. The epidermal layer above the split appeared to be normal. The basement membrane was intact, along with normal underlying connective tissue. Discussion: Management consisted of decompression of blisters followed by topical antibiotics. Oral corticosteroids were given for control of blistering, since they reduce collagenase activity. Avoidance of trauma is essential aspect of management: baby was nursed with care to prevent occurrence of new lesions. Therapy and counseling sessions were scheduled for the parents.

Keywords: Bart Syndrome. Epidermolysis Bullosa Dystrophica. Transient bullous dermolysis of the newborn. Oral manifestations.
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Introduction

Bart Syndrome or congenital transient mechanobullous dermatosis is one of the lesser known presentations of epidermolysis bullosa (EB) (1). It is a rare inherited skin blistering disorder. It is an autosomal dominant inheritance, caused by mutation in the collagen type VII gene on chromosome 3p. It is characterized by congenital skin defects with bullae of the extremities and intertriginous areas, congenital localized absence of skin anterior aspect of lower extremities and dorsa of the feet, erosions of mouth and dystrophic nails. All of these features are remitted within a few years (2).

Case report

A female newborn aged 3 days old was admitted to the neonatal ward of Dermatology Department, Adesh Institute of Medical Sciences and Research, Bathinda, Punjab, India, with chief complaint of erosions of the legs and difficulty in feeding. Hence an oral diagnostician was called upon.

A detailed case history revealed positive family history; maternal uncle had history of EB. On examination it was observed that eroded lesions were distributed over the hands, feet, and chest and on the face over the cheeks bilaterally. The skin lesions showed denuded areas with bullae rupturing easily to reveal painful eroded areas. The entire medial aspect of the left leg showed raw eroded skin extending from the knee to the toes (Figure 1). The ruptured bullae were also distributed over the chest, face and hands (Figure 2). The nails showed dystrophic changes. The lips were erythematous, eroded with tissue tags (Figure 3). On intraoral examination, the entire labial mucosa showed eroded, crustated lesions. Intra oral blisters were seen over anterior palate which subsequently ruptured to form erosions.

Based on the above findings, provisional diagnosis of EB was given. Routine blood investigations revealed raised erythrocyte sedimentation rate (ESR). A blister was induced by rubbing normal looking...
corticosteroids were given for control of blistering, since they reduce collagenase activity. Therapy and counselling sessions were scheduled for the parents. Genetic counselling is an essential part of management of families in whom there is an individual with EB. Skin care instructions were as follows: the baby was nursed on a foam mattress one inch in thickness. New bullae were drained by puncturing them with a sterile needle and roof left in situ. The baby was bathed in warm water containing a little potassium permanganate. Vaseline gauge was applied to the eroded areas. Avoidance of trauma is essential aspect of management, baby nursed with care to prevent occurrence of new lesions.

Discussion

The term ‘epidermolysis bullosa’ was first coined by Koebner. Hallopeau was the first to distinguish between simple (non-scarring) and dystrophic (scarring) forms of the disease. Bart Syndrome was originally described in a large family in 1966 by Bart (3).

Management

The patient was given dressing over the left leg, along with topical antibiotic cream. Oral skin for one minute. The skin biopsy was taken from the area after 10 min. Histopathological examination revealed split localized to the epidermis. The epidermal layer above the split appeared to be normal. The basement membrane was intact, along with normal underlying connective tissue, suggestive of EB (Figure 4).

The above clinical findings of bullae of skin, oral mucosa and dystrophic nails and histopathological findings helped arrive at the final diagnosis of Bart Syndrome.
Bart Syndrome is a triad of bullous involvement of cutaneous, oral lesions and dystrophic nails. The onset is at birth or in early infancy. Mechanical trauma experienced during delivery may lead to bullae and erosions at the sites of trauma. The common sites are hand, feet and other areas of friction including oral mucosa. Blistering is usually worse in summer and in absence of secondary infection, heals without scarring. Oral ulcers formed as a result of rupture of bullae heal without remarkable sequelae, may develop in the newborn due to vigorous sucking (4).

Though its etiology is unknown, it is believed to be an autoimmune disorder. The antigen is type VII collagen, a 290 KD glycoprotein which is specifically found in basement membranes beneath stratified squamous epithelia (5-7). Physical trauma in utero has also been proposed as a mechanism to explain denuded areas of the limbs (2).

A detailed history should include (8):

1) mapping of the family pedigree;
2) evidence of consanguinity;
3) age of onset;
4) seasonal effects;
5) fever: shows extent of disease activity;
6) cutaneous findings.

Differential diagnosis can be Bullous pemphigoid, Bullous lichen planus, Cicatricial pemphigoid (9). Histopathological examination of the skin biopsy localizes the split to the epidermis (EB simplex) or the basement membrane zone (junctional and dystrophic EB). Bart Syndrome can present as simplex, junctional or dystrophic forms. Electron microscopy can differentiate between EB simplex (intraepidermal split), junctional EB (intraamina lucida) and dystrophic EB (sub lamina lucida) (10) Specific antibodies to antigen, type VII collagen can be found by immunofluorescence techniques (11). The form of EB in this case was simplex. Only histopathologic examination by light microscopy was done. EB simplex has onset in childhood or in adult life. Lesions are exacerbated in warm weather. These lesions usually heal without scarring and secondary infection is a common complication. Prognosis of EB simplex is favourable. Oral lesions occur immediately after first feed over palate, tongue and labial mucosa in junctional EB. The bullae rupture to form widespread erosions.

Gastrointestinal tract involvement shows blistering from oesophagus to large intestine leading to pyloric stenosis. This impairs calorie intake causing anaemia and growth retardation. Laryngeal oedema causes asphyxia. Prognosis is grave. Dystrophic EB is characterised by generalized blistering and erosions which heal with atrophy, scarring and milia formation. Squamous cell carcinoma occasionally develops at the site of chronic trauma. Prognosis is unfavourable (4).

Management of this syndrome consists of prevention of trauma, a cool environment and decompression of blisters followed by topical antibiotics.

a) In neonates blisters are punctured with a sterile needle, bathed in warm water containing a little amount of potassium permanganate. Vaseline gauge is applied to eroded areas. If infected, an antibiotic cream is used.
b) In older infants and toddlers erosions are covered with topical antiseptics such as 10% povidine iodine or 1% silver sulfadiazine cream (4).

Systemic treatment modalities include oral corticosteroids in high doses for control of blistering in both dystrophic and junctional EB. They reduce collagenase activity of fibroblasts. Retinoic acid inhibits collagenase production. 13-cis-retinoic acid at a dose of 0.4 mg/kg/day reduces the number of lesions. A daily dose of 300-1000 IU of Vitamin E diminishes blister formation (12). Phenytoin can be used in recurrent and junctional EB at a dose 100-300 mg once daily for 6-8 weeks (13, 14). This decreases at least 50% of bullous lesions. It acts by decreasing synthesis of collagenase (15).

Partial thickness skin autografts may be used for reconstructive surgery in neonates affected by severe forms of EB (4).

Conclusions

Oral professionals have their role in the diagnosis and management of Bart Syndrome. Recognizing the oral clinical features and the prompt management is important.
References


