A Case of Non-Hallopeau-Siemens Recessive Dystrophic Epidermolysis Bullosa

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Dystrophic epidermolysis bullosa (DEB) is a rare group of heritable mechanobullous disorders that are characterized by blistering and scarring of the skin and mucosae and these lesions are induced by minor trauma. DEB is also associated with nail dystrophy. DEB can be inherited either in an autosomal recessive or dominant fashion. Regardless of the mode of inheritance, DEB is caused by defects of the ultrastructural entity known as the anchoring fibril, which results in separation of the sublamina densa. Recessive DEB (RDEB) is classified into Hallopeau-Siemens and non-Hallopeau-Siemens. We herein report on a case of non-Hallopeau-Siemens RDEB and there was no family history of this malady, and we present the clinical, histological and electron microscopy findings.

INTRODUCTION

Dystrophic epidermolysis bullosa (DEB) is a group of heritable mechanobullous disorders that’s characterized by blistering and scarring of the skin and mucosae and nail dystrophy that’s all induced by minor trauma. The ultrastructural hallmark of DEB is tissue separation at the sublamina densa level, and this condition is usually associated with an abnormal quality or quantity of anchoring fibrils at the dermal-epidermal junction. Dystrophic epidermolysis bullosa is inherited either in an autosomal dominant or recessive mode. Recessive DEB (RDEB) is a much more serious form than dominant DEB, with the blistering generally being present from birth. RDEB is classified into Hallopeau-Siemens and non-Hallopeau-Siemens. We describe here a case of non-Hallopeau-Siemens RDEB that was diagnosed based on the family history and the clinical, histological and electron microscopic findings.
Fig. 1. (A) Bullae and erosions on the lower legs and feet. (B) Skin erosions and wound exudate on the dorsum of the left hand and fingers.

Fig. 2. Subepidermal blister formation and flattened rete ridges are observed (H&E, ×100).

Fig. 3. Electron microscopy showing the presence lamina densa (arrowhead) attached to epidermal keratinocytes at the roof of the cleft. Nu: nucleus of keratinocyte, bar=1.0 μm (×15,000).

dermal keratinocytes at the roof of the cleft, and this was compatible with the diagnosis of DEB (Fig. 3). Based on the family history and the clinical, histological and electron microscopic findings, this case was diagnosed as non-Hallopeau-Siemens recessive dystrophic epidermolysis bullosa. She was treated with regular skin cleaning, dressings and 2% mupirocin ointment. She was also treated with phenytoin (7 mg/d), vitamin E (150 mg/d) and systemic antibiotics (amikacin 50 mg/d, vancomycin 240 mg/d) for 15 days, but no improvement was observed. She was scheduled to undergo genetic studies, but she was transferred to another tertiary hospital by her parent’s wishes.

DISCUSSION

Epidermolysis bullosa (EB) is a complex group of inherited or acquired cutaneous diseases in which bullous lesions arise after normal levels of physical trauma. EB can be divided into three major general categories, based on the level of tissue separation within the cutaneous basement membrane zone that separates the epidermis from the dermis. In epidermolysis bullosa simplex (EBS), blisters occur within the epidermis. In junctional epidermolysis bullosa (JEB), the separation is in the lamina lucida of the dermoepidermal junction and in DEB, the lesions arise in the upper dermis. The mode of inheritance is typically autosomal dominant in EBS and autosomal recessive in JEB. DEB can be either autosomal dominant or autosomal recessive. RDEB is a much more serious form than dominant DEB, with blistering generally present from birth. RDEB is divided into Hallopeau-Siemens and non-Hallopeau-Siemens. RDEB is the most severe form of DEB and it is responsible for widespread mucocutaneous blistering leading to fusion.
of the digits, nail loss, flexural contractures, esophageal strictures, narrowing of the trachea or larynx as well as oral and ocular erosions. Malnutrition, anemia and growth retardation commonly occur. Anal and genitourinary involvements may also be present. Squamous cell carcinoma is a common complication of the cutaneous scarring and it is a significant cause of mortality. All the patients with RDEB who lack the cutaneous and extracutaneous features so characteristic of the Hallopeau-Siemens subtype are included under the term non-Hallopeau-Siemens RDEB.

According to a National Epidermolysis Bullosa Registry report, 50 EB cases occur per 1 million live births. Of these cases, approximately 92% are EBS, 5% are DEB, 1% is JEB and 2% are unclassified. DEB occurs in all racial and ethnic groups and it equally affects males and females. The epidemiologic data on this illness in Korea has been insufficient. In 1993, Rho et al. surveyed 79 patients diagnosed with EB from 1970 to 1992 at the departments of dermatology and pediatrics in university hospitals and dermatologic departments in general hospitals in Korea. The epidemiologic study showed 8 cases of RDEB, and Kim et al. reported 3 more cases in 2001. Yet because of the low incidence of EB in Korea, solid statistical data on this disease is still lacking.

The diagnosis of DEB is based on the clinical signs, the histopathology, the electron microscopy and, if available, genetic studies. Separation of the epidermis and dermis was found on the histologic biopsy study of this case, and the electron microscopy showed the lamina densa attached to epidermal keratinocytes at the roof of the cleft, which is compatible with the diagnosis of DEB. Blisters and erosions were seen at birth, and this is also common in RDEB. The symptoms were generalized, but any extracutaneous features seen in the Hallopeau-Siemens subtype were not present, and the cutaneous features were mild. For those reasons, we were able to diagnose the patient as non-Hallopeau-Siemens RDEB.

Treatment for this disease is mainly symptomatic and supportive, as the currently available therapies cannot correct the underlying molecular defect. The treatment regimen is tailored to the severity and extent of the skin and systemic involvement and this usually entails a combination of wound management, infection control, surgical management as needed and nutritional support. Administration of phenytoin has been experimented with for some recessive DEB cases for its inhibitory action on collagenase and it has been suggested as a therapeutic modality, but several studies have reportedly various responses. Our patient was given phenytoin with vitamin E and systemic antibiotics, but she failed to show a significant response.

The symptoms of dominant DEB are mostly minor and it is possible to lead a normal life. But RDEB may lead to death that’s caused by sepsis, depending on the degree of invasion, and the complication rates for such maladies as esophageal strictures and skin cancer are higher. Therefore, a multidisciplinary approach is needed for these patients to assure an optimal outcome. Especially, counseling and advise for the details of skin care must be provided and then reinforced to the parents of children who suffer with RDEB.

REFERENCES


11. Abahussein AA, al-Zayir AA, Mostafa WZ, Okoro AN.