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Ectodermal Dysplasia with Bilateral Punctal Agenesis in a Nigerian Child: A Case Report

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Summary

Ectodermal dysplasia (ED) is a hereditary condition that occurs worldwide. It is due to abnormality in the development of skin and its appendages (hair, teeth and nail). ED also presents with ocular manifestations. This case of ED is reported to highlight ocular manifestations such as bilateral punctal agenesis, madarosis and blepharitis. The need for a multi-disciplinary approach in managing such cases is underscored.

Keywords: Ectodermal Dysplasia, Nigeria, Punctal agenesis, Skin appendages.

Introduction

Ectodermal dysplasia (ED) is a heterogeneous group of congenital disorders, mostly inherited in an X-linked pattern, first reported by Thurnam in 1848.¹ The term “ectodermal dysplasia” was coined by Weech in 1929.² ED consists of a large, heterogeneous, congenital, non-progressive group of disorders defined by primary defects in the development of two or more tissues derived embryologically from the ectoderm. The primary tissues affected are the teeth, skin and its appendages (sebaceous glands, eccrine glands, hair follicles and nails).³

Over 170 specific syndromes have been described with different modes of inheritance.⁴ ED classification system was first proposed by Freire-Maia and Pinheiro in 1982,⁵ with an additional modification in 1994 and 2001.⁶ The patients were divided into subgroups based on the presence or absence of trichодysplasia, abnormal dentition, onchodysplasia and dyshidrosis. The most common syndromes within this group are hypohidrotic ED and hidrotic ED.⁷ The hypohidrotic and hidrotic forms have varying manifestations of hair, teeth and nail abnormalities. The hypohidrotic form (Christ-Siemens-Touraine syndrome) is the most common, with significant involvement of the sweat glands, and mostly inherited as an X-linked recessive disorder compared to the hidrotic form (Clouston syndrome), which spares the sweat glands and has an autosomal dominant mode of inheritance.⁷ Many cases with ocular manifestations of ED have been reported worldwide.⁸-¹² Familusi et al. reported the first ED case in Nigeria in 1975. Following this, many cases of ED have subsequently been reported in Nigeria, but no single case has been reported on ophthalmic manifestations in Nigerian subjects.
Worldwide, the frequency of the different forms of ED in a given population varies greatly and ranges from 1 in 10,000 to as low as 1 in 100,000 births. Any tissue embryologically derived from the ectoderm, be it surface ectoderm or neuroectoderm, could be affected. Some cases may even have a mesodermal component to this disorder. [22]

Therefore, this case of hidrotic ectodermal dysplasia with bilateral punctal agenesis, madarosis and blepharitis in a Nigerian child is reported. This case deserves reporting because of its unusual features and difficulty in providing optimal management.

**Case Description**

A 4-year-old female child was brought to the Department of Ophthalmology, University of Ilorin Teaching Hospital, Ilorin, Nigeria, by the parents with a history of bilateral tearing since birth. There were associated recurrent redness, itching of the eyelid margin and purulent discharge which subsides with the instillation of topical antibiotics (chloramphenicol ointment and ofloxacin eye drop). There was no history of reduction in vision, swelling in the medial canthi region, heat intolerance, dysphonia, hoarseness of voice or hearing defect. The mother gave a history of discoloration and irregular growth of the child's dentitions and nails as well as uneven growth and loss of scalp hair. There was no family history of similar abnormalities or consanguineous marriage.

Physical examination revealed frontal bossing with sparse, fine hair and loss of hair on the scalp along the central part of the sagittal suture (Figure 1), nail dystrophy involving all digits (Figure 2) and malformed teeth (Figure 3).

Ocular examination of both eyes revealed visual acuity of 6/9 in the Right Eye and Left Eye respectively, blepharitis, madarosis (Figure 4) and bilateral absence of the puncta in the upper or lower lids (all four puncta) (Figure 5) with the presence of lacrimal papillae on examination under anaesthesia.

The posterior segment in both eyes was essentially normal, and intra-ocular pressure was within the normal range bilaterally. Based on the history and examination findings, ectodermal dysplasia was diagnosed. All the requested investigations (Full Blood Count, Random Blood Glucose, and urinalysis) gave results within the normal range. The parents were extensively counselled on the child's clinical condition. The child was co-managed with the dermatologist, paediatrician and dental surgeon (prosthodontist). She subsequently had bilateral conjunctivodacryocystorhinostomy (C-DCR) with the placement of a Lester Jones tube in both eyes, and blepharitis was adequately managed with topical antibiotics (chloramphenicol ointment, and gutfloxacin), gutf diclofenac and scrubbing of the eyelids with baby shampoo.

The researchers obtained informed consent to use the child's data for this report from the child's mother. Ethical approval for the study was also obtained from the Research Ethics Committee of the Federal Medical Centre, Birnin-Kebbi, Kebbi State, Nigeria.

**Discussion**

The index case is similar to the hidrotic form as there was no history of heat intolerance or recurrent fever, even though there was no family history of ED. Several ED syndromes may also manifest in association with midfacial defects, mainly cleft lip and palate. [4] ED is often seen among whites [23] but rare among blacks. Hidrotic ED has been reported in several members of a French-Canadian family. [3] Clinical recognition of ED varies from birth to childhood, depending on the severity of symptoms and the recognition of associated complications, as many patients are not diagnosed until infancy or childhood when...
dental, nail, and hair abnormalities have started manifesting. [3]

As highlighted above, patients with ED have several systemic manifestations, but of particular interest in this report are the ocular abnormalities. The previously reported ophthalmologic manifestations include strabismus, telecanthus, fused lids at birth, blepharophimosis, entropion, absence of eyelashes, bilateral eyelid cysts, agenesis of lacrimal puncta, dacycystitis, blepharitis, conjunctivitis, deficient meibomian gland function, and corneal limbal deficiency. [8-12]

There are few reports of the anhidrotic form among Nigerians. [24-26] However, to the authors' knowledge, this is the first case of the hidrotic form with ocular manifestations in Nigeria. ED is a disorder well documented in the dental and dermatological literature but in very few ophthalmological literature. The index child had the typical ocular associations, including complete agenesis of the upper and lower puncta bilaterally, madarosis and blepharitis, which have been documented in previous studies. [8-12]

The previous studies on ED with ocular association described bilateral involvement of the lacrimal apparatus, [27-29] also present in the index case. This informed our decision to do a conjunctivo-dacryocysto-rhinostomy (C-DCR) with a Lester Jones tube placement. The procedure was successful but post-operatively, the child developed a bilateral infection of the tubes, necessitating the removal of the tubes. Previous studies had attributed the tube infection to the stagnation of tears due to the poorly developed sino-nasal structures and chronic rhino-sinusitis, which are part of the disease spectrum. [28] However, the index child had no detectable disease manifestation on the ocular surface and posterior segment. Nevertheless, the child was closely followed up for early detection of other lately manifesting abnormalities in the eye, as documented by Marshall et al. [29]

This case report emphasizes the various pattern of systemic and ocular manifestations of the disease. Managing these patients, especially
those with significant systemic abnormalities can be challenging and requires a multi-disciplinary approach. The managing team should include the ophthalmologist, dental surgeon, plastic surgeon, dermatologist, and speech therapist. The family and caregivers must be very supportive as the disorder can lead to low self-esteem and reduced quality of life, considering the patient’s obvious teeth, skin, hair and nail abnormalities.

Conclusion

Ectodermal dysplasia remains a rare condition seen in our environment. Management requires a multi-disciplinary approach. Therefore, it is essential to create awareness of this rare disease.

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